ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

ARICLAIM 40 mg hard gastro-resistant capsules.

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each capsule contains 40 mg of duloxetine (as hydrochloride).

Excipients: sucrose 11.5 mg.

For a full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard gastro-resistant capsule.

Opaque orange body, imprinted with '40mg' and an opaque blue cap, imprinted with '9545'.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

ARICLAIM is indicated for women for the treatment of moderate to severe Stress Urinary Incontinence (SUI), (see section 5.1).

Treatment of diabetic peripheral neuropathic pain in adults.

4.2 Posology and method of administration

Stress Urinary Incontinence:

The recommended dose of ARICLAIM is 40 mg twice daily without regard to meals. After 2-4 weeks of treatment, patients should be re-assessed in order to evaluate the benefit and tolerability of the therapy. Some patients may benefit from starting treatment at a dose of 20 mg twice daily for two weeks before increasing to the recommended dose of 40 mg twice daily. Dose escalation may decrease, though not eliminate, the risk of nausea and dizziness.

However, limited data are available to support the efficacy of ARICLAIM 20 mg twice daily. A 20 mg capsule is also available.

The efficacy of ARICLAIM has not been evaluated for longer than 3 months in placebo-controlled studies. The benefit of treatment should be re-assessed at regular intervals.

Combining ARICLAIM with a pelvic floor muscle training (PFMT) program may be more effective than either treatment alone. It is recommended that consideration be given to concomitant PFMT.

Diabetic Peripheral Neuropathic Pain:

The starting and recommended maintenance dose is 60 mg daily with or without food. Dosages above 60 mg once daily, up to a maximum dose of 120 mg per day administered in evenly divided doses, have been evaluated from a safety perspective in clinical trials. The plasma concentration of duloxetine displays large inter-individual variability (see 5.2). Hence, some patients that respond insufficiently to 60 mg may benefit from a higher dose.

Response to treatment should be evaluated after 2 months. In patients with inadequate initial response, additional response after this time is unlikely.

The therapeutic benefit should be reassessed regularly (at least every three months) (see Section 5.1).

Hepatic insufficiency:

ARICLAIM should not be used in patients with liver disease resulting in hepatic impairment (see section 4.3).

Renal insufficiency:

No dosage adjustment is necessary for patients with mild or moderate renal dysfunction (creatinine clearance 30 to 80 ml/min). ARICLAIM should not be used in patients with severe renal impairment (creatinine clearance <30 ml/min; see section 4.3).

Elderly

No dosage adjustment is recommended for elderly patients solely on the basis of age. Caution should be exercised when treating the elderly

Children and adolescents:

There is no experience in children (see section 4.4).

Discontinuation of treatment:

Abrupt discontinuation should be avoided. When stopping treatment with ARICLAIM the dose should be gradually reduced over a period of at least one to two weeks in order to reduce the risk of withdrawal reactions (see sections 4.4 and 4.8). If intolerable symptoms occur following a decrease in the dose or upon discontinuation of treatment, then resuming the previously prescribed dose may be considered. Subsequently, the physician may continue decreasing the dose, but at a more gradual rate.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients.

Liver disease resulting in hepatic impairment (see section 5.2).

ARICLAIM should not be used in combination with nonselective, irreversible Monoamine Oxidase Inhibitors - MAOIs (see section 4.5).

ARICLAIM should not be used in combination with CYP1A2 inhibitors, like fluvoxamine, ciprofloxacin or enoxacine since the combination results in elevated plasma concentrations of duloxetine (see section 4.5).

Severe renal impairment (creatinine clearance <30 ml/min) (see section 4.4).

The initiation of treatment with ARICLAIM is contraindicated in patients with uncontrolled hypertension that could expose patients to a potential risk of hypertensive crisis (see sections 4.4 and 4.8).

4.4 Special warnings and precautions for use

Mania and Seizures

ARICLAIM should be used with caution in patients with a history of mania or a diagnosis of bipolar disorder, and/or seizures.

Use with antidepressants

The use of ARICLAIM in combination with antidepressants (especially with SSRI, SNRI and reversible MAOIs) is not recommended (see below "Depression, suicidal ideation and behaviour" and Section 4.5).

St John's wort

Undesirable effects may be more common during concomitant use of ARICLAIM and herbal preparations containing St John's wort (Hypericum perforatum).

Mydriasis

Mydriasis has been reported in association with duloxetine, therefore, caution should be used when prescribing duloxetine in patients with increased intraocular pressure, or those at risk of acute narrowangle glaucoma.

Blood pressure and heart rate

Duloxetine has been associated with an increase in blood pressure and clinically significant hypertension in some patients. This may be due to the noradrenergic effect of duloxetine. Cases of hypertensive crisis have been reported with duloxetine, especially in patients with pre-existing hypertension. Therefore,in patients with known hypertension and/or other cardiac disease, blood pressure monitoring is recommended, especially during the first month of treatment. Duloxetine should be used with caution in patients whose conditions could be compromised by an increased heart rate or by an increase in blood pressure. Caution should also be exercised when duloxetine is used with medicinal products that may impair its metabolism (see section 4.5). For patients who experience a sustained increase in blood pressure while receiving duloxetine either dose reduction or gradual discontinuation should be considered (see section 4.8). In patients with uncontrolled hypertension duloxetine should not be initiated (see section 4.3).

Renal impairment

Increased plasma concentrations of duloxetine occur in patients with severe renal impairment on haemodialysis (creatinine clearance <30 ml/min). For patients with severe renal impairment, see section 4.3. See section 4.2 for information on patients with mild or moderate renal dysfunction.

Haemorrhage

There have been reports of bleeding abnormalities, such as ecchymoses, purpura and gastrointestinal haemorrhage with selective serotonin reuptake inhibitors (SSRIs) and serotonin/noradrenaline reuptake inhibitors (SNRIs). Caution is advised in patients taking anticoagulants and/or medicinal products known to affect platelet function, and in patients with known bleeding tendencies.

Discontinuation of treatment

Withdrawal symptoms when treatment is discontinued are common, particularly if discontinuation is abrupt (see section 4.8). In a clinical trial adverse events seen on abrupt treatment discontinuation occurred in approximately 44% of patients treated with ARICLAIM and 24% of patients taking placebo.

The risk of withdrawal symptoms seen with SSRI's and SNRI's may be dependent on several factors including the duration and dose of therapy and the rate of dose reduction. The most commonly reported reactions are listed in section 4.8. Generally the symptoms are mild to moderate, however, in some patients they may be severe in intensity. They usually occur within the first few days of discontinuing treatment, but there have been very rare reports of such symptoms in patients who have inadvertently missed a dose. Generally these symptoms are self-limiting and usually resolve within 2 weeks, though in some individuals they may be prolonged (2-3 months or more). It is therefore advised that duloxetine should be gradually tapered when discontinuing treatment over a period of no less than 2 weeks, according to the patient's needs (see section 4.2).

Hyponatremia

Hyponatremia has been reported rarely, predominantly in the elderly, when administering ARICLAIM. Caution is required in patients at increased risk for hyponatremia; such as elderly, cirrhotic, or dehydrated patients or patients treated with diuretics. Hyponatremia may be due to a syndrome of inappropriate anti-diuretic hormone secretion (SIADH).

Depression, suicidal ideation and behaviour

Although ARICLAIM is not indicated for the treatment of depression, its active ingredient (duloxetine) also exists as an antidepressant medication. Depression is associated with an increased risk of suicidal thoughts, self harm and suicide (suicide-related events). This risk persists until significant remission occurs. As improvement may not occur during the first few weeks or more of treatment, patients should be closely monitored until such improvement occurs. It is general clinical experience that the risk of suicide may increase in the early stages of recovery. Patients with a history

of suicide-related events or those exhibiting a significant degree of suicidal thoughts prior to commencement of treatment are known to be at a greater risk of suicidal thoughts or suicidal behaviour, and should receive careful monitoring during treatment. A meta-analysis of placebo-controlled clinical trials of antidepressant medicinal products in psychiatric disorders showed an increased risk of suicidal behaviour with antidepressants compared to placebo in patients less than 25 years old

Cases of suicidal thoughts and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.8). Physicians should encourage patients to report any distressing thoughts or feelings or depressive symptoms at any time. If while on ARICLAIM therapy, the patient develops agitation or depressive symptoms, specialised medical advice should be sought, as depression is a serious medical condition. If a decision to initiate antidepressant pharmacological therapy is taken, the gradual discontinuation of ARICLAIM is recommended (see Section 4.2).

Use in children and adolescents under 18 years of age

No clinical trials have been conducted with duloxetine in paediatric populations. ARICLAIM should not be used in the treatment of children and adolescents under the age of 18 years. Suicide-related behaviours (suicide attempts and suicidal thoughts), and hostility (predominantly aggression, oppositional behaviour and anger), were more frequently observed in clinical trials among children and adolescents treated with antidepressants compared to those treated with placebo. Long-term safety data in children and adolescents concerning growth, maturation and cognitive and behavioural development are lacking.

Medicinal products containing duloxetine

Duloxetine is used under different trademarks in several indications (treatment of diabetic neuropathic pain, major depressive episodes as well as stress urinary incontinence). The use of more than one of these products concomitantly should be avoided.

Hepatitis/increased liver enzymes

Cases of liver injury, including severe elevations of liver enzymes (>10 times upper limit of normal), hepatitis and jaundice have been reported with duloxetine (see section 4.8). Most of them occurred during the first months of treatment. The pattern of liver damage was predominantly hepatocellular. Duloxetine should be used with caution in patients treated with other medicinal products associated with hepatic injury.

Akathisia/psychomotor restlessness

The use of duloxetine has been associated with the development of akathisia, characterised by a subjectively unpleasant or distressing restlessness and need to move often accompanied by an inability to sit or stand still. This is most likely to occur within the first few weeks of treatment. In patients who develop these symptoms, increasing the dose may be detrimental.

Sucrose

ARICLAIM gastro-resistant capsules contain sucrose. Patients with rare hereditary problems of fructose intolerance, glucose-galactose malabsorption or sucrose-isomaltase insufficiency should not take this medicine.

4.5 Interaction with other medicinal products and other forms of interaction

Monoamine oxidase inhibitors (MAOIs): due to the risk of serotonin syndrome, ARICLAIM should not be used in combination with nonselective, irreversible monoamine oxidase inhibitors (MAOIs), or within at least 14 days of discontinuing treatment with an MAOI. Based on the half-life of duloxetine, at least 5 days should be allowed after stopping ARICLAIM before starting an MAOI (see section 4.3).

Serotonin syndrome: in rare cases, serotonin syndrome has been reported in patients using SSRIs (e.g. paroxetine, fluoxetine) concomitantly with serotonergic medicinal products. Caution is advisable if ARICLAIM is used concomitantly with serotonergic antidepressants like SSRIs, tricyclics like

clomipramine or amitriptyline, St John's wort (Hypericum perforatum), venlafaxine or triptans, tramadol, pethidine and tryptophan.

CNS medicinal products: caution is advised when ARICLAIM is taken in combination with other centrally acting medicinal products or substances, including alcohol and sedative medicinal products (benzodiazepines, morphinomimetics, antipsychotics, phenobarbital, sedative antihistamines).

Effect of duloxetine on other medicinal products

Medicinal products metabolised by CYP1A2: The pharmacokinetics of theophylline, a CYP1A2 substrate, were not significantly affected by co-administration with duloxetine (60 mg twice daily). Medicinal products metabolised by CYP2D6: Duloxetine is a moderate inhibitor of CYP2D6. When duloxetine was administered at a dose of 60 mg twice daily with a single dose of desipramine, a CYP2D6 substrate, the AUC of desipramine increased 3-fold. The co-administration of duloxetine (40 mg twice daily) increases steady state AUC of tolterodine (2 mg twice daily) by 71 %, but does not affect the pharmacokinetics of its active 5-hydroxyl metabolite and no dosage adjustment is recommended. Caution is advised if ARICLAIM is co-administered with medicinal products that are predominantly metabolised by CYP2D6 (risperidone, tricyclic antidepressants [TCAs] such as nortriptyline, amitriptyline, and imipramine) particularly if they have a narrow therapeutic index (such as flecainide, propafenone and metoprolol).

Oral contraceptives and other steroidal agents: results of *in vitro* studies demonstrate that duloxetine does not induce the catalytic activity of CYP3A. Specific *in vivo* drug interaction studies have not been performed.

Anticoagulants and antiplatelet agents: Caution should be exercised when duloxetine is combined with oral anticoagulants or antiplatelet agents due to a potential increased risk of bleeding. Furthermore, increases in INR values have been reported when duloxetine was co-administered with warfarin.

Effects of other medicinal products on duloxetine

Antacids and H2 antagonists: co-administration of ARICLAIM with aluminium- and magnesium-containing antacids or with famotidine had no significant effect on the rate or extent of duloxetine absorption after administration of a 40 mg oral dose.

Inhibitors of CYP1A2: because CYP1A2 is involved in duloxetine metabolism, concomitant use of ARICLAIM with potent inhibitors of CYP1A2 is likely to result in higher concentrations of duloxetine. Fluvoxamine (100 mg once daily), a potent inhibitor of CYP1A2, decreased the apparent plasma clearance of duloxetine by about 77% and increased AUC_{0-t} 6-fold. Therefore ARICLAIM should not be administered in combination with potent inhibitors of CYP1A2 like fluvoxamine (see section 4.3).

Inducers of CYP1A2: Population pharmacokinetic studies analyses have shown that smokers have almost 50% lower plasma concentrations of duloxetine compared with non-smokers.

4.6 Pregnancy and lactation

Pregnancy

There are no data on the use of duloxetine in pregnant women. Studies in animals have shown reproductive toxicity at systemic exposure levels (AUC) of duloxetine lower than the maximum clinical exposure (see section 5.3). The potential risk for humans is unknown. As with other serotonergic medicinal product, discontinuation symptoms may occur in the neonate after maternal duloxetine use near term. ARICLAIM should be used in pregnancy only if the potential benefit justifies the potential risk to the foetus. Women should be advised to notify their physician if they become pregnant, or intend to become pregnant, during therapy.

Breast feeding

Duloxetine is very weakly excreted into human milk based on a study of 6 lactating patients, who did not breast feed their children . The estimated daily infant dose on a mg/kg basis is approximately 0.14% of the maternal dose (see section 5.2). As the safety of duloxetine in infants is not known, the use of ARICLAIM while breast-feeding is not recommended.

4.7 Effects on ability to drive and use machines

No studies on the effects on the ability to drive and use machines have been performed. ARICLAIM may be associated with sedation and dizziness. Patients should be instructed that if they experience sedation or dizziness they should avoid potentially hazardous tasks such as driving or operating machinery.

4.8 Undesirable effects

Stress Urinary Incontinence:

Table 1 gives the adverse reactions observed from spontaneous reporting and in placebo-controlled clinical trials (comprising a total of 8241 patients, 4504 on duloxetine and 3737 on placebo) in SUI and other lower urinary tract disorders.

The most commonly reported adverse events in patients treated with ARICLAIM in clinical trials in SUI and other lower urinary tract disorders were nausea, dry mouth fatigue and constipation. The data analysis of four 12-week, placebo-controlled clinical trials in patients with SUI, including 958 duloxetine-treated and 955 placebo-treated patients, showed that the onset of the reported adverse events typically occurred in the first week of therapy. However, the majority of the most frequent adverse events were mild to moderate and resolved within 30 days of occurrence (e.g. nausea).

Table 1: Adverse reactions

Frequency are defined as: Very common ($\geq 1/10$), common ($\geq 1/100$ to <1/10), uncommon ($\geq 1/1,000$ to <1/100), rare ($\geq 1/10,000$ and <1/1,000), very rare (<1/10,000), not known (cannot be estimated from the available data).

Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Investigations		<u>.</u>		•	•
		Weight decrease Weight increase Blood cholesterol increased Creatine phosphokinase increased			
Cardiac Disorder.	5	1		T	1
		Palpitations Tachycardia			Supra- ventricular arrhythmia, mainly atrial fibrillation
Nervous System D	isorders				
	Headache Dizziness Tremor Lethargy Somnolence	Poor quality sleep Disturbance in attention Nervousness	Dyskinesia Myoclonus Restless legs syndrome		Serotonin syndrome Psychomotor restlessness Convulsions ¹

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
	Paraesthesia	Dysgeusia			Akathisia Extrapyramida-l symptoms
Eye Disorders		•			
	Blurred vision	Visual disturbance Mydriasis	Glaucoma		
Ear and Labyrinth	Disorders	1vi y di lasis			
Der eine Bue yr inn	Vertigo	Tinnitus ¹ Ear pain			
Respiratory, thora	cic and mediastina	l disorders	<u> </u>		•
		Yawning	Epistaxis Throat tightness		
Gastrointestinal D	isorders				•
Nausea (22.8%) Dry mouth (12.1%) Constipation (10.3%)	Diarrhoea Vomiting Dyspepsia	Gastroenteritis Stomatitis Gastritis Flatulence Eructation Breath odour	Haematochezia		Gastrointestina-l haemorrhage
Renal and Urinary	y Disorders				
		Urinary hesitation Dysuria Nocturia Urine odour abnormal	Urine flow decreased Polyuria		Urinary retention
Skin and Subcutan	eous Tissue Disord	lers	1		1
	Sweating increased	Rash Increased tendency to bruise Night sweats Cold sweat Dermatitis contact Urticaria	Photo-sensitivity reactions		Stevens-Johnson Syndrome Angioneurotic oedema
Musculoskeletal a	nd connective tissu		I I		1
		Muscle spasm Muscle tightness Musculo- skeletal pain Trismus	Muscle twitching		
Endocrine disorde	ers	Hyma	<u> </u>		
		Hypo- thyroidism			
Metabolism and N	utrition Disorders				<u> </u>
	Appetite decreased	Dehydration	Hyperglycemia (reported especially in diabetic patients) Hyponatremia		SIADH
Infections and infe	estations				

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
		Laryngitis			
Vascular Disorde	rs		<u>. </u>		
	Flushing	Syncope ² Blood pressure increase	Hypertensive crisis Orthostatic hypotension ² Peri pheral coldness		Hypertension
Ganaral Disordar	rs and Administration	on Sita Conditions			
Fatigue (10.9%)	Abdominal pain Asthenia Chills	Malaise Feeling abnormal Feeling cold Feeling hot Thirst	Gait disturbance		Chest pain
Immune system di	isorders		I		
		Hyper- sensitivity disorder	Anaphylactic reaction		
Hepato-biliary di	sorders				
		Hepatitis ³ Elevated liver enzymes (ALT, AST, alkaline phosphatase) Acute liver injury			Hepatic failure Jaundice
Reproductive Syst	tem and Breast Disc		<u>. </u>		
		Menopausal symptoms Gynaecologica-l haemorrhage			
Psychiatric Disor	ders		ı L		1
	Insomnia Anxiety Sleep disorder Agitation Libido decreased	Disorientation Abnormal dreams Apathy BruxismOrgasm abnormal	Hallucinations reported after treatn		Suicidal behaviour Suicidal ideation ⁴ Mania Aggression and anger ⁵

¹ Cases of convulsion and cases of tinnitus have also been reported after treatment discontinuation.

Discontinuation of duloxetine (particularly when abrupt) commonly leads to withdrawal symptoms. Dizziness, sensory disturbances (including paraesthesia), sleep disturbances (including insomnia and intense dreams), agitation or anxiety, nausea and/or vomiting, tremor, headache, irritability, diarrhoea, hyperhydrosis and vertigo are the most commonly reported reactions.

²Cases of orthostatic hypotension and syncope have been reported especially at the initiation of treatment

³See section 4.4

⁴Cases of suicidal ideation and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.4)

⁵Cases of aggression and anger have been reported particularly early in treatment of after treatment discontinuation.

Generally, for SSRIs and SNRIs, these events are mild to moderate and self-limiting, however, in some patients they may be severe and/or prolonged. It is therefore advised that when duloxetine treatment is no longer required, gradual discontinuation by dose tapering should be carried out (see sections 4.2 and 4.4).

Electrocardiograms were obtained from 755 duloxetine-treated patients with SUI and 779 placebotreated patients in 12-week clinical trials. The heart rate-corrected QT interval in duloxetine-treated patients did not differ from that seen in placebo-treated patients.

In the 12 week acute phase of three clinical trials of duloxetine in patients with diabetic neuropathic pain, small but statistically significant increases in fasting blood glucose were observed in duloxetine-treated patients . HbA $_{1c}$ was stable in both duloxetine-treated and placebo-treated patients. In the extension phase of these studies, which lasted up to 52 weeks, there was an increase in HbA $_{1c}$ in both the duloxetine and routine care groups, but the mean increase was 0.3% greater in the duloxetine-treated group. There was also a small increase in fasting blood glucose and in total cholesterol in duloxetine-treated patients while those laboratory tests showed a slight decrease in the routine care group.

Diabetic Peripheral Neuropathic Pain

Table 2 gives the adverse reactions observed from spontaneous reporting and in placebo-controlled clinical trials (comprising a total of 6828 patients, 4199 on duloxetine and 2629 on placebo).

The most commonly observed adverse reactions in patients with diabetic neuropathic pain treated with ARICLAIM were nausea, headache, dry mouth somnolence and dizziness. However, the majority of common adverse reactions were mild to moderate, they usually started early in therapy, and most tended to subside even as therapy was continued.

Table 2: Adverse reactions

Frequency estimate: Very common ($\geq 1/10$), common ($\geq 1/100$ and <1/10), uncommon ($\geq 1/1,000$ and <1/100), rare ($\geq 1/10,000$ and <1/1,000), very rare (<1/10,000), not known (cannot be estimated from the available data).

Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Investigations					
	Weight decrease	Weight increase	Blood		
		Creatine	cholesterol		
		phosphokinase	increased		
		increased			
Cardiac Disorder	rs				
	Palpitations	Tachycardia			
		Supra-			
		ventricular			
		arrhythmia,			
		mainly atrial			
		fibrillation			
Nervous System I	Disorders				
Headache	Tremor	Myoclonus			Serotonin
(14.3%)	Paraesthesia	NervousnessDis	Convulsion ¹		syndrome
Somnolence		turbance in			Extra-pyramidal
(10.7%)		attention			symptoms
Dizziness		Lethargy			Akathisia
(10.2%)		Dysgeusia			Psychomotor
		Dyskinesia			restlessness

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
		Restless legs syndrome Poor quality sleep			
Eye Disorders					
	Blurred vision	Mydriasis Visual disturbance	Glaucoma		
Ear and Labyrint	h Disorders	<u> </u>	<u> </u>		
	Tinnitus ¹	Vertigo Ear pain			
Respiratory, thore	acic and mediastin		T		
	Yawning	Throat tightness Epistaxis			
Gastrointestinal I		Gastroenteritis	Stomatitis		Castus intestina 1
Nausea (24.3%) Dry mouth (12.8%)	Constipation Diarrhoea Vomiting Dyspepsia Flatulence	Eructation Gastritis	Breath odour Haematochezi-a		Gastrointestina-l haemorrhage
Renal and Urinar		1			
		Urinary Retention Dysuria Urinary hesitation Nocturia Polyuria Urine flow decreased	Urine odour abnormal		
Skin and Subcuta	neous Tissue Disor		T		T
	Sweating increased Rash	Night sweats Urticaria Dermatitis contact Cold sweat Photosensitivity reactions Increased tendency to bruise			Angio-neurotic oedema Stevens- Johnson Syndrome
Musculoskeletal (and connective tissi	ue disorders			
	Musculo- skeletal pain Muscle tightness Muscle spasm	Muscle twitching	Trismus		
Endocrine disord	ers	1	T **		_
			Нуро-		

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
			thyroidism		
Metabolism and I	Nutrition Disorders	T	1		<u> </u>
	Decreased Appetite	Hyperglycemi-a (reported especially in diabetic patients)	Dehydration Hyponatremia		SIADH
Infections and inf	estations	,			
		Laryngitis			
Vascular Disorde		T :			T .
	Flushing	Blood pressure increase Peripheral coldness Orthostatic hypotension ² Syncope ²			Hypertension Hypertensive crisis
General Disorder	rs and Administratio	on Site Conditions			
	Fatigue Abdominal pain	Feeling abnormal Feeling cold Thirst Chills Malaise Feeling hot Gait disturbance			Chest pain
Immune system di	isorders				
			Hyper- sensitivity disorder Anaphylactic reaction		
Hepato-biliary di	sorders	T	T		T
		Elevated liver enzymes (ALT, AST, alkaline phosphatase) Hepatitis ³ Acute liver injury			Jaundice Hepatic failure
Reproductive Syst	l tem and Breast Disc	orders	<u> </u>		
- sp. samoure syst	Erectile dysfunction	Ejaculation disorder Ejaculation delayed Sexual dysfunction Gynaecological haemorrhage	Menopausal symptoms		

Very common	Common	Uncommon	Rare	Very Rare	Frequency not				
					known				
Psychiatric Disor	Psychiatric Disorders								
	Insomnia	Sleep disorder	Mania		Suicidal				
	Agitation	Bruxism	Hallucinations		ideation ⁵				
	Libido	Disorientation	Aggression and		Suicidal ⁵				
	decreased	Apathy	anger ⁴		behaviour				
	Anxiety								
	Orgasm								
	abnormal								
	Abnormal								
	dreams								

¹Cases of convulsion and cases of tinnitus have also been reported after treatment discontinuation.

Discontinuation of duloxetine (particularly when abrupt) commonly leads to withdrawal symptoms. Dizziness, sensory disturbances (including paraesthesia), sleep disturbances (including insomnia and intense dreams), agitation or anxiety, nausea and/or vomiting, tremor, headache, irritability, diarrhoea, hyperhydrosis and vertigo are the most commonly reported reactions.

Generally, for SSRIs and SNRIs, these events are mild to moderate and self-limiting, however, in some patients they may be severe and/or prolonged. It is therefore advised that when duloxetine treatment is no longer required, gradual discontinuation by dose tapering should be carried out (see sections 4.2 and 4.4).

In the 12 week acute phase of three clinical trials of duloxetine in patients with diabetic neuropathic pain, small but statistically significant increases in fasting blood glucose were observed in duloxetine-treated patients. HbA1c was stable in both duloxetine-treated and placebo-treated patients. In the extension phase of these studies, which lasted up to 52 weeks, there was an increase in HbA1c in both the duloxetine and routine care groups, but the mean increase was 0.3% greater in the duloxetine-treated group. There was also a small increase in fasting blood glucose and in total cholesterol in duloxetine-treated patients while those laboratory tests showed a slight decrease in the routine care group.

Electrocardiograms were obtained from 528 duloxetine-treated and 205 placebo-treated patients with diabetic neuropathic pain in clinical trials lasting up to 13-weeks. The heart rate-corrected QT interval in duloxetine-treated patients did not differ from that seen in placebo-treated patients. No clinically significant differences were observed for QT, PR, QRS, or QTcB measurements between duloxetine-treated and placebo-treated patients.

4.9 Overdose

Cases of overdoses, alone or in combination with other medicinal products, with duloxetine doses of 5400mg were reported. Some fatalities have occurred, primarily with mixed overdoses, but also with duloxetine alone at a dose of approximately 1000 mg. Signs and symptoms of overdose (duloxetine alone or in combination with other medicinal products) included somnolence, coma, serotonin syndrome, seizures, vomiting and tachycardia.

No specific antidote for duloxetine is known but if serotonin syndrome ensues, specific treatment (such as with cyproheptadine and/or temperature control) may be considered. A free airway should be established. Monitoring of cardiac and vital signs is recommended, along with appropriate symptomatic and supportive measures. Gastric lavage may be indicated if performed soon after

²Cases of orthostatic hypotension and syncope have been reported especially at the initiation of treatment.

³See section 4.4

⁴Cases of aggression and anger have been reported particularly early in treatment of after treatment discontinuation.

⁵Cases of suicidal ideation and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.4)

ingestion or in symptomatic patients. Activated charcoal may be useful in limiting absorption. Duloxetine has a large volume of distribution and forced diuresis, haemoperfusion, and exchange perfusion are unlikely to be beneficial.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other antidepressants. ATC code: N06AX21

Duloxetine is a combined serotonin (5-HT) and noradrenaline (NA) reuptake inhibitor. It weakly inhibits dopamine reuptake with no significant affinity for histaminergic, dopaminergic, cholinergic and adrenergic receptors.

In animal studies, increased levels of 5-HT and NE in the sacral spinal cord, lead to increased urethral tone via enhanced pudendal nerve stimulation to the urethral striated sphincter muscle only during the storage phase of the micturition cycle. A similar mechanism in women is believed to result in stronger urethral closure during urine storage with physical stress that could explain the efficacy of duloxetine in the treatment of women with SUI.

Duloxetine normalised pain thresholds in several preclinical models of neuropathic and inflammatory pain and attenuated pain behaviour in a model of persistent pain. The pain inhibitory action of duloxetine is believed to be a result of potentiation of descending inhibitory pain pathways within the central nervous system.

Stress Urinary Incontinence: The efficacy of duloxetine 40 mg given twice daily in the treatment of SUI was established in four double-blind, placebo-controlled studies, that randomised 1913 women (22 to 83 years) with SUI; of these, 958 patients were randomised to duloxetine and 955 to placebo. The primary efficacy measures were Incontinence Episode Frequency (IEF) from diaries and an incontinence specific quality of life questionnaire score (I-QOL).

Incontinence Episode Frequency: in all four studies the duloxetine-treated group had a 50% or greater median decrease in IEF compared with 33% in the placebo-treated group. Differences were observed at each visit after 4 weeks (duloxetine 54%, and placebo 22%), 8 weeks (52% and 29%), and 12 weeks (52% and 33%) of medication.

In an additional study limited to patients with severe SUI, all responses with duloxetine were achieved within 2 weeks.

The efficacy of ARICLAIM has not been evaluated for longer than 3 months in placebo-controlled studies. The clinical benefit of ARICLAIM compared with placebo has not been demonstrated in women with mild SUI, defined in randomised trials as those with IEF < 14 per week. In these women, ARICLAIM may provide no benefit beyond that afforded by more conservative behavioural interventions.

Quality of Life: Incontinence Quality of Life (I-QOL) questionnaire scores were significantly improved in the duloxetine-treated patient group compared with the placebo-treated group (9.2 versus 5.9 score improvement, p<.001). Using a global improvement scale (PGI), significantly more women using duloxetine considered their symptoms of stress incontinence to be improved with treatment compared with women using placebo (64.6% versus 50.1%, p<.001).

ARICLAIM and Prior Continence Surgery: there are limited data that suggest that the benefits of ARICLAIM are not diminished in women with stress urinary incontinence who have previously undergone continence surgery.

ARICLAIM and Pelvic Floor Muscle Training (PFMT): during a 12-week blinded, randomised, controlled study, ARICLAIM demonstrated greater reductions in IEF compared with either placebo treatment or with PFMT alone. Combined therapy (duloxetine + PFMT) showed greater improvement in both pad use and condition-specific quality of life measures than ARICLAIM alone or PFMT alone.

Diabetic Peripheral Neuropathic Pain:

The efficacy of duloxetine as a treatment for diabetic neuropathic pain was established in 2 randomised, 12-week, double-blind, placebo-controlled, fixed dose studies in adults (22 to 88 years) having diabetic neuropathic pain for at least 6 months. Patients meeting diagnostic criteria for major depressive disorder were excluded from these trials. The primary outcome measure was the weekly mean of 24-hour average pain, which was collected in a daily diary by patients on an 11-point Likert scale.

In both studies, duloxetine 60 mg once daily and 60 mg twice daily significantly reduced pain compared with placebo. The effect in some patients was apparent in the first week of treatment. The difference in mean improvement between the two active treatment arms was not significant. At least 30% reported pain reduction was recorded in approximately 65% of duloxetine treated patients versus 40% for placebo. The corresponding figures for at least 50% pain reduction were 50% and 26% respectively. Clinical response rates (50% or greater improvement in pain) were analysed according to whether or not the patient experienced somnolence during treatment. For patients not experiencing somnolence, clinical response was observed in 47% of patients receiving duloxetine and 27% patients on placebo. Clinical response rates in patients experiencing somnolence were 60% on duloxetine and 30% on placebo. Patients not demonstrating a pain reduction of 30% within 60 days of treatment were unlikely to reach this level during further treatment.

In an open label long-term uncontrolled study, the pain reduction in patients responding to 8-weeks of acute treatment of ARICLAIM 60 mg once daily was maintained for a further 6-months as measured by change on the Brief Pain Inventory(BPI) 24-hour average pain item.

5.2 Pharmacokinetic properties

Duloxetine is administered as a single enantiomer. Duloxetine is extensively metabolised by oxidative enzymes (CYP1A2 and the polymorphic CYP2D6), followed by conjugation. The pharmacokinetics of duloxetine demonstrate large intersubject variability (generally 50-60%), partly due to gender, age, smoking status and CYP2D6 metaboliser status.

Duloxetine is well absorbed after oral administration with a C_{max} occurring 6 hours post dose. The absolute oral bioavailability of duloxetine ranged from 32% to 80% (mean of 50%; N=8 subjects). Food delays the time to reach the peak concentration from 6 to 10 hours and it marginally decreases the extent of absorption (approximately 11%).

Duloxetine is approximately 96% bound to human plasma proteins. Duloxetine binds to both albumin and alpha-1 acid glycoprotein. Protein binding is not affected by renal or hepatic impairment.

Duloxetine is extensively metabolised and the metabolites are excreted principally in urine. Both CYP2D6 and CYP1A2 catalyse the formation of the two major metabolites glucuronide conjugate of 4-hydroxy duloxetine and sulphate conjugate of 5-hydroxy,6-methoxy duloxetine. Based upon *in vitro* studies, the circulating metabolites of duloxetine are considered pharmacologically inactive. The pharmacokinetics of duloxetine in patients who are poor metabolisers with respect to CYP2D6 has not been specifically investigated. Limited data suggest that the plasma levels of duloxetine are higher in these patients.

The elimination half-life of duloxetine after an oral dose ranges from 8 to 17 hours (mean of 12 hours). After an intravenous dose the plasma clearance of duloxetine ranges from 22 l/hr to 46 l/hr (mean of 36 l/hr) After an oral dose the apparent plasma clearance of duloxetine ranges from 33 to 261 l/hr (mean 101 l/hr).

Special populations:

Gender: pharmacokinetic differences have been identified between males and females (apparent plasma clearance is approximatively 50% lower in females). Based upon the overlap in the range of clearance, gender-based pharmacokinetic differences do not justify the recommendation for using a lower dose for female patients.

Age: pharmacokinetic differences have been identified between younger and elderly females (\geq 65 years) (AUC increases by about 25% and half-life is about 25% longer in the elderly), although the magnitude of these changes is not sufficient to justify adjustments to the dose. As a general recommendation, caution should be exercised when treating the elderly (see sections 4.2 and 4.4).

Renal impairment: end stage renal disease (ESRD) patients receiving dialysis had a 2-fold higher duloxetine C_{max} and AUC values compared to healthy subjects. Pharmacokinetic data on duloxetine is limited in patients with mild or moderate renal impairment.

Hepatic insufficiency: moderate liver disease (Child Pugh Class B) affected the pharmacokinetics of duloxetine. Compared with healthy subjects, the apparent plasma clearance of duloxetine was 79% lower, the apparent terminal half-life was 2.3 times longer, and the AUC was 3.7 times higher in patients with moderate liver disease. The pharmacokinetics of duloxetine and its metabolites have not been studied in patients with mild or severe hepatic insufficiency.

Nursing mothers: The disposition of duloxetine was studied in 6 lactating women who were at least 12-weeks postpartum. Duloxetine is detected in breast milk, and steady-state concentrations in breast milk are about one-fourth those in plasma. The amount of duloxetine in breast milk is approximately 7 μg/day while on 40 mg twice daily dosing. Lactation did not influence duloxetine pharmacokinetics.

5.3 Preclinical safety data

Duloxetine was not genotoxic in a standard battery of tests and was not carcinogenic in rats. Multinucleated cells were seen in the liver in the absence of other histopathological changes in the rat carcinogenicity study. The underlying mechanism and the clinical relevance are unknown.

Female mice receiving duloxetine for 2 years had an increased incidence of hepatocellular adenomas and carcinomas at the high dose only (144 mg/kg/day), but these were considered to be secondary to hepatic microsomal enzyme induction. The relevance of this mouse data to humans is unknown. Female rats receiving duloxetine before and during mating and early pregnancy had a decrease in maternal food consumption and body weight, oestrous cycle disruption, decreased live birth indices and progeny survival, and progeny growth retardation at systemic exposure levels estimated to be at the most at maximum clinical exposure (AUC). In an embryotoxicity study in the rabbit, a higher incidence of cardiovascular and skeletal malformations was observed at systemic exposure levels below the maximum clinical exposure (AUC). No malformations were observed in another study testing a higher dose of a different salt of duloxetine. In pre/postnatal toxicity study in the rat, duloxetine induced adverse behavioural effects in the offspring at systemic exposures levels below maximum clinical exposure (AUC).

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Capsule content:

Hypromellose. Hypromellose acetate succinate Sucrose Sugar spheres Talc Titanium dioxide (E171) Triethyl citrate.

Capsule shell:

Gelatin Sodium Lauryl Sulfate Titanium Dioxide (E171) Indigo Carmine (E132) Red Iron oxide (E172) Yellow Iron Oxide (E172) Edible black ink.

Edible Ink: Black Iron Oxide-Synthetic (E172) Propylene glycol Shellac.

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years.

6.4 Special precautions for storage

Store in the original package in order to protect from moisture. Do not store above 30°C.

6.5 Nature and contents of container

Polyvinylchloride (PVC), Polyethylene (PE), and Polychlorotrifluoroethylene (PCTFE) blister sealed with an aluminium foil.

Packs of 28, 56, 98, 140 and 196 (2x98) capsules.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

Eli Lilly Nederland BV, Grootslag 1-5, NL-3991 RA Houten, The Netherlands.

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/04/283/002 EU/1/04/283/003

EU/1/04/283/004

EU/1/04/283/005

EU/1/04/283/006

9.	DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION
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11 August 2004

10. DATE OF REVISION OF THE TEXT

1. NAME OF THE MEDICINAL PRODUCT

ARICLAIM 20 mg hard gastro-resistant capsules.

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each capsule contains 20 mg of duloxetine (as hydrochloride).

Excipients: sucrose 5.7 mg.

For a full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard gastro-resistant capsule.

Opaque blue body, imprinted with '20 mg' and an opaque blue cap, imprinted with '9544'.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

ARICLAIM is indicated for women for the treatment of moderate to severe Stress Urinary Incontinence (SUI), (see section 5.1).

Treatment of diabetic peripheral neuropathic pain in adults.

4.2 Posology and method of administration

Stress Urinary Incontinence:

The recommended dose of ARICLAIM is 40 mg twice daily without regard to meals. After 2-4 weeks of treatment, patients should be re-assessed in order to evaluate the benefit and tolerability of the therapy. Some patients may benefit from starting treatment at a dose of 20 mg twice daily for two weeks before increasing to the recommended dose of 40 mg twice daily. Dose escalation may decrease, though not eliminate, the risk of nausea and dizziness.

However, limited data are available to support the efficacy of ARICLAIM 20 mg twice daily.

The efficacy of ARICLAIM has not been evaluated for longer than 3 months in placebo-controlled studies. The benefit of treatment should be re-assessed at regular intervals.

Combining ARICLAIM with a pelvic floor muscle training (PFMT) program may be more effective than either treatment alone. It is recommended that consideration be given to concomitant PFMT.

Diabetic Peripheral Neuropathic Pain:

The starting and recommended maintenance dose is 60 mg daily with or without food. Dosages above 60 mg once daily, up to a maximum dose of 120 mg per day administered in evenly divided doses, have been evaluated from a safety perspective in clinical trials. The plasma concentration of duloxetine displays large inter-individual variability (see 5.2). Hence, some patients that respond insufficiently to 60 mg may benefit from a higher dose.

Response to treatment should be evaluated after 2 months. In patients with inadequate initial response, additional response after this time is unlikely.

The therapeutic benefit should be reassessed regularly (at least every three months) (see Section 5.1).

Hepatic insufficiency:

ARICLAIM should not be used in patients with liver disease resulting in hepatic impairment (see section 4.3).

Renal insufficiency:

No dosage adjustment is necessary for patients with mild or moderate renal dysfunction (creatinine clearance 30 to 80 ml/min). ARICLAIM should not be used in patients with severe renal impairment (creatinine clearance <30 ml/min; see section 4.3).

Elderly:

No dosage adjustment is recommended for elderly patients solely on the basis of age. Caution should be exercised when treating the elderly.

Children and adolescents:

There is no experience in children (see section 4.4).

Discontinuation of treatment:

Abrupt discontinuation should be avoided. When stopping treatment with ARICLAIM the dose should be gradually reduced over a period of at least one to two weeks in order to reduce the risk of withdrawal reactions (see sections 4.4 and 4.8). If intolerable symptoms occur following a decrease in the dose or upon discontinuation of treatment, then resuming the previously prescribed dose may be considered. Subsequently, the physician may continue decreasing the dose, but at a more gradual rate.).

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients.

Liver disease resulting in hepatic impairment (see section 5.2).

ARICLAIM should not be used in combination with nonselective, irreversible Monoamine Oxidase Inhibitors - MAOIs (see section 4.5).

ARICLAIM should not be used in combination with CYP1A2 inhibitors, like fluvoxamine, ciprofloxacin or enoxacine since the combination results in elevated plasma concentrations of duloxetine (see section 4.5).

Severe renal impairment (creatinine clearance <30 ml/min) (see section 4.4).

The initiation of treatment with ARICLAIM is contraindicated in patients with uncontrolled hypertension that could expose patients to a potential risk of hypertensive crisis (see sections 4.4 and 4.8).

4.4 Special warnings and precautions for use

Mania and Seizures

ARICLAIM should be used with caution in patients with a history of mania or a diagnosis of bipolar disorder, and/or seizures.

Use with antidepressants

The use of ARICLAIM in combination with antidepressants (especially with SSRI, SNRI and reversible MAOIs) is not recommended (see below "Depression, suicidal ideation and behaviour" and Section 4.5).

St John's wort

Undesirable effects may be more common during concomitant use of ARICLAIM and herbal preparations containing St John's wort (Hypericum perforatum).

Mvdriasis

Mydriasis has been reported in association with duloxetine, therefore, caution should be used when prescribing duloxetine in patients with increased intraocular pressure, or those at risk of acute narrowangle glaucoma.

Blood pressure and heart rate

Duloxetine has been associated with an increase in blood pressure and clinically significant hypertension in some patients. This may be due to the noradrenergic effect of duloxetine. Cases of hypertensive crisis have been reported with duloxetine, especially in patients with pre-existing hypertension. Therefore,in patients with known hypertension and/or other cardiac disease, blood pressure monitoring is recommended, especially during the first month of treatment. Duloxetine should be used with caution in patients whose conditions could be compromised by an increased heart rate or by an increase in blood pressure. Caution should also be exercised when duloxetine is used with medicinal products that may impair its metabolism (see section 4.5). For patients who experience a sustained increase in blood pressure while receiving duloxetine either dose reduction or gradual discontinuation should be considered (see section 4.8). In patients with uncontrolled hypertension duloxetine should not be initiated (see section 4.3).

Renal impairment

Increased plasma concentrations of duloxetine occur in patients with severe renal impairment on haemodialysis (creatinine clearance <30 ml/min). For patients with severe renal impairment, see section 4.3. See section 4.2 for information on patients with mild or moderate renal dysfunction.

Haemorrhage

There have been reports of bleeding abnormalities, such as ecchymoses, purpura and gastrointestinal haemorrhage with selective serotonin reuptake inhibitors (SSRIs) and serotonin/noradrenaline reuptake inhibitors (SNRIs). Caution is advised in patients taking anticoagulants and/or medicinal products known to affect platelet function, and in patients with known bleeding tendencies.

Discontinuation of treatment

Withdrawal symptoms when treatment is discontinued are common, particularly if discontinuation is abrupt (see section 4.8). In a clinical trial, adverse events seen on abrupt treatment discontinuation occurred in approximately 44% of patients treated with ARICLAIM and 24% of patients taking placebo.

The risk of withdrawal symptoms seen with SSRI's and SNRI's may be dependent on several factors including the duration and dose of therapy and the rate of dose reduction. The most commonly reported reactions are listed in section 4.8. Generally the symptoms are mild to moderate, however, in some patients they may be severe in intensity. They usually occur within the first few days of discontinuing treatment, but there have been very rare reports of such symptoms in patients who have inadvertently missed a dose. Generally these symptoms are self-limiting and usually resolve within 2 weeks, though in some individuals they may be prolonged (2-3 months or more). It is therefore advised that duloxetine should be gradually tapered when discontinuing treatment over a period of no less than 2 weeks, according to the patient's needs (see section 4.2).

Hyponatremia

Hyponatremia has been reported rarely, predominantly in the elderly, when administering ARICLAIM. Caution is required in patients at increased risk for hyponatremia; such as elderly, cirrhotic, or dehydrated patients or patients treated with diuretics. Hyponatremia may be due to a syndrome of inappropriate anti-diuretic hormone secretion (SIADH).

Depression, suicidal ideation and behaviour

Although ARICLAIM is not indicated for the treatment of depression, its active ingredient (duloxetine) also exists as an antidepressant medication. Depression is associated with an increased risk of suicidal thoughts, self harm and suicide (suicide-related events). This risk persists until significant remission occurs. As improvement may not occur during the first few weeks or more of treatment, patients should be closely monitored until such improvement occurs. It is general clinical

experience that the risk of suicide may increase in the early stages of recovery. Patients with a history of suicide-related events or those exhibiting a significant degree of suicidal thoughts prior to commencement of treatment are known to be at a greater risk of suicidal thoughts or suicidal behaviour, and should receive careful monitoring during treatment. A meta-analysis of placebo-controlled clinical trials of antidepressant medicinal products in psychiatric disorders showed an increased risk of suicidal behaviour with antidepressants compared to placebo in patients less than 25 years old

Cases of suicidal thoughts and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.8). Physicians should encourage patients to report any distressing thoughts or feelings or depressive symptoms at any time. If while on ARICLAIM therapy, the patient develops agitation or depressive symptoms, specialised medical advice should be sought, as depression is a serious medical condition. If a decision to initiate antidepressant pharmacological therapy is taken, the gradual discontinuation of ARICLAIM is recommended (see Section 4.2).

Use in children and adolescents under 18 years of age

No clinical trials have been conducted with duloxetine in paediatric populations. ARICLAIM should not be used in the treatment of children and adolescents under the age of 18 years. Suicide-related behaviours (suicide attempts and suicidal thoughts), and hostility (predominantly aggression, oppositional behaviour and anger), were more frequently observed in clinical trials among children and adolescents treated with antidepressants compared to those treated with placebo. Long-term safety data in children and adolescents concerning growth, maturation and cognitive and behavioural development are lacking.

Medicinal products containing duloxetine

Duloxetine is used under different trademarks in several indications (treatment of diabetic neuropathic pain, major depressive episodes as well as stress urinary incontinence). The use of more than one of these products concomitantly should be avoided.

Hepatitis/increased liver enzymes

Cases of liver injury, including severe elevations of liver enzymes (>10 times upper limit of normal), hepatitis and jaundice have been reported with duloxetine (see section 4.8). Most of them occurred during the first months of treatment. The pattern of liver damage was predominantly hepatocellular. Duloxetine should be used with caution in patients treated with other medicinal products associated with hepatic injury.

Akathisia/psychomotor restlessness

The use of duloxetine has been associated with the development of akathisia, characterised by a subjectively unpleasant or distressing restlessness and need to move often accompanied by an inability to sit or stand still. This is most likely to occur within the first few weeks of treatment. In patients who develop these symptoms, increasing the dose may be detrimental.

Sucrose

ARICLAIM gastro-resistant capsules contain sucrose. Patients with rare hereditary problems of fructose intolerance, glucose-galactose malabsorption or sucrose-isomaltase insufficiency should not take this medicine.

4.5 Interaction with other medicinal products and other forms of interaction

Monoamine oxidase inhibitors (MAOIs): due to the risk of serotonin syndrome, ARICLAIM should not be used in combination with nonselective, irreversible monoamine oxidase inhibitors (MAOIs), or within at least 14 days of discontinuing treatment with an MAOI. Based on the half-life of duloxetine, at least 5 days should be allowed after stopping ARICLAIM before starting an MAOI (see section 4.3).

Serotonin syndrome: in rare cases, serotonin syndrome has been reported in patients using SSRIs (e.g. paroxetine, fluoxetine) concomitantly with serotonergic medicinal products. Caution is advisable if

ARICLAIM is used concomitantly with serotonergic antidepressants like SSRIs, tricyclics like clomipramine or amitriptyline, St John's wort (Hypericum perforatum), venlafaxine or triptans, tramadol, pethidine and tryptophan.

CNS medicinal products: caution is advised when ARICLAIM is taken in combination with other centrally acting medicinal products or substances, including alcohol and sedative medicinal products (benzodiazepines, morphinomimetics, antipsychotics, phenobarbital, sedative antihistamines).

Effect of duloxetine on other medicinal products

Medicinal products metabolised by CYP1A2: The pharmacokinetics of theophylline, a CYP1A2 substrate, were not significantly affected by co-administration with duloxetine (60 mg twice daily). Medicinal products metabolised by CYP2D6: Duloxetine is a moderate inhibitor of CYP2D6. When duloxetine was administered at a dose of 60 mg twice daily with a single dose of desipramine, a CYP2D6 substrate, the AUC of desipramine increased 3-fold. The co-administration of duloxetine (40 mg twice daily) increases steady state AUC of tolterodine (2 mg twice daily) by 71 %, but does not affect the pharmacokinetics of its active 5-hydroxyl metabolite and no dosage adjustment is recommended. Caution is advised if ARICLAIM is co-administered with medicinal products that are predominantly metabolised by CYP2D6 (risperidone, tricyclic antidepressants [TCAs] such as nortriptyline, amitriptyline, and imipramine) particularly if they have a narrow therapeutic index (such as flecainide, propafenone and metoprolol).

Oral contraceptives and other steroidal agents: results of *in vitro* studies demonstrate that duloxetine does not induce the catalytic activity of CYP3A. Specific *in vivo* drug interaction studies have not been performed.

Anticoagulants and antiplatelet agents: Caution should be exercised when duloxetine is combined with oral anticoagulants or antiplatelet agents due to a potential increased risk of bleeding. Furthermore, increases in INR values have been reported when duloxetine was co-administered with warfarin.

Effects of other medicinal products on duloxetine

Antacids and H2 antagonists: co-administration of ARICLAIM with aluminium- and magnesium-containing antacids or with famotidine had no significant effect on the rate or extent of duloxetine absorption after administration of a 40 mg oral dose.

Inhibitors of CYP1A2: because CYP1A2 is involved in duloxetine metabolism, concomitant use of ARICLAIM with potent inhibitors of CYP1A2 is likely to result in higher concentrations of duloxetine. Fluvoxamine (100 mg once daily), a potent inhibitor of CYP1A2, decreased the apparent plasma clearance of duloxetine by about 77% and increased AUC_{0-t}6-fold. Therefore ARICLAIM should not be administered in combination with potent inhibitors of CYP1A2 like fluvoxamine (see section 4.3).

Inducers of CYP1A2: Population pharmacokinetic studies analyses have shown that smokers have almost 50% lower plasma concentrations of duloxetine compared with non-smokers.

4.6 Pregnancy and lactation

Pregnancy

There are no data on the use of duloxetine in pregnant women. Studies in animals have shown reproductive toxicity at systemic exposure levels (AUC) of duloxetine lower than the maximum clinical exposure (see section 5.3). The potential risk for humans is unknown.

As with other serotonergic medicinal product, discontinuation symptoms may occur in the neonate after maternal duloxetine use near term. ARICLAIM should be used in pregnancy only if the potential benefit justifies the potential risk to the foetus. Women should be advised to notify their physician if they become pregnant, or intend to become pregnant, during therapy.

Breast feeding

Duloxetine is very weakly excreted into human milk based on a study of 6 lactating patients, who did not breast feed their children . The estimated daily infant dose on a mg/kg basis is approximately 0.14% of the maternal dose (see section 5.2). As the safety of duloxetine in infants is not known, the use of ARICLAIM while breast-feeding is not recommended.

4.7 Effects on ability to drive and use machines

No studies on the effects on the ability to drive and use machines have been performed. ARICLAIM may be associated with sedation and dizziness. Patients should be instructed that if they experience sedation or dizziness they should avoid potentially hazardous tasks such as driving or operating machinery.

4.8 Undesirable effects

Stress Urinary Incontinence:

Table 1 gives the adverse reactions observed from spontaneous reporting and in placebo-controlled clinical trials (comprising a total of 8241 patients, 4504 on duloxetine and 3737 on placebo) in SUI and other lower urinary tract disorders.

The most commonly reported adverse events in patients treated with ARICLAIM in clinical trials in SUI and other lower urinary tract disorders were nausea, dry mouth fatigue and constipation. The data analysis of four 12-week, placebo-controlled clinical trials in patients with SUI, including 958 duloxetine-treated and 955 placebo-treated patients, showed that the onset of the reported adverse events typically occurred in the first week of therapy. However, the majority of the most frequent adverse events were mild to moderate and resolved within 30 days of occurrence (e.g. nausea).

Table 1: Adverse reactions

Frequency are defined as: Very common ($\geq 1/10$), common ($\geq 1/100$ to <1/10), uncommon ($\geq 1/1,000$ to <1/100), rare ($\geq 1/10,000$ and <1/1,000), very rare (<1/10,000), not known (cannot be estimated from the available data).

Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Investigations					
		Weight decrease Weight increase Blood cholesterol increased Creatine phosphokinase increased			
Cardiac Disorders	5				
		Palpitations Tachycardia			Supra- ventricular arrhythmia, mainly atrial fibrillation
Nervous System D	isorders				
	Headache Dizziness Tremor Lethargy Somnolence	Poor quality sleep Disturbance in attention Nervousness	Dyskinesia Myoclonus Restless legs syndrome		Serotonin syndrome Psychomotor restlessness Convulsions ¹

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
	Paraesthesia	Dysgeusia			Akathisia Extrapyramida-l symptoms
Eye Disorders					
	Blurred vision	Visual disturbance Mydriasis	Glaucoma		
Ear and Labyrinth	n Disorders				•
	Vertigo	Tinnitus ¹ Ear pain			
Respiratory, thora	icic and mediastina	ıl disorders	-		
		Yawning	Epistaxis Throat tightness		
Gastrointestinal L	Disorders				
Nausea (22.8%) Dry mouth (12.1%) Constipation (10.3%)	Diarrhoea Vomiting Dyspepsia	Gastroenteritis Stomatitis Gastritis Flatulence Eructation Breath odour	Haematochezia		Gastrointestina-l haemorrhage
Renal and Urinar	y Disorders				
		Urinary hesitation Dysuria Nocturia Urine odour abnormal	Urine flow decreased Polyuria		Urinary retention
Skin and Subcutar	ieous Tissue Disord	ders	1		1
	Sweating increased	Rash Increased tendency to bruise Night sweats Cold sweat Dermatitis contact Urticaria	Photo-sensitivity reactions		Stevens-Johnson Syndrome Angioneurotic oedema
Musculoskeletal a	nd connective tissu		<u> </u>		
		Muscle spasm Muscle tightness Musculo- skeletal pain Trismus	Muscle twitching		
Endocrine disorde	ers	1	<u>l</u>		l
		Hypo- thyroidism			
Metabolism and N	lutrition Disorders		<u> </u>		'
	Appetite decreased	Dehydration	Hyperglycemia (reported especially in diabetic patients) Hyponatremia		SIADH
Infections and infe	estations	1	113 ponutioniu		

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
		Laryngitis			
Vascular Disorde			<u></u>		
	Flushing	Syncope ² Blood pressure increase	Hypertensive crisis Orthostatic hypotension ² Peri pheral coldness		Hypertension
General Disorder	s and Administratio	n Site Conditions			
Fatigue (10.9%)	Abdominal pain Asthenia Chills	Malaise Feeling abnormal Feeling cold Feeling hot Thirst	Gait disturbance		Chest pain
Immune system di	sorders	<u> </u>	L		_
		Hyper- sensitivity disorder	Anaphylactic reaction		
Hepato-biliary dis	sorders	•	l l		1
		Hepatitis ³ Elevated liver enzymes (ALT, AST, alkaline phosphatase) Acute liver injury			Hepatic failure Jaundice
Reproductive Syst	em and Breast Disc	, , , , , , , , , , , , , , , , , , ,	I.		L
		Menopausal symptoms Gynaecologica-l haemorrhage			
Psychiatric Disord		Diagniant stiss	Hallmainetiene		Cuinida!
	Insomnia Anxiety Sleep disorder Agitation Libido decreased	Disorientation Abnormal dreams Apathy Bruxism Orgasm abnormal	Hallucinations		Suicidal behaviour Suicidal ideation ⁴ Mania Aggression and anger ⁵

¹ Cases of convulsion and cases of tinnitus have also been reported after treatment discontinuation.

Discontinuation of duloxetine (particularly when abrupt) commonly leads to withdrawal symptoms. Dizziness, sensory disturbances (including paraesthesia), sleep disturbances (including insomnia and

²Cases of orthostatic hypotension and syncope have been reported especially at the initiation of treatment

³See section 4.4

⁴Cases of suicidal ideation and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.4)

⁵Cases of aggression and anger have been reported particularly early in treatment of after treatment discontinuation.

intense dreams), agitation or anxiety, nausea and/or vomiting, tremor, headache, irritability, diarrhoea, hyperhydrosis and vertigo are the most commonly reported reactions.

Generally, for SSRIs and SNRIs, these events are mild to moderate and self-limiting, however, in some patients they may be severe and/or prolonged. It is therefore advised that when duloxetine treatment is no longer required, gradual discontinuation by dose tapering should be carried out (see sections 4.2 and 4.4).

Electrocardiograms were obtained from 755 duloxetine-treated patients with SUI and 779 placebotreated patients in 12-week clinical trials. The heart rate-corrected QT interval in duloxetine-treated patients did not differ from that seen in placebo-treated patients.

In the 12 week acute phase of three clinical trials of duloxetine in patients with diabetic neuropathic pain, small but statistically significant increases in fasting blood glucose were observed in duloxetine-treated patients . HbA $_{\rm lc}$ was stable in both duloxetine-treated and placebo-treated patients. In the extension phase of these studies, which lasted up to 52 weeks, there was an increase in HbA $_{\rm lc}$ in both the duloxetine and routine care groups, but the mean increase was 0.3% greater in the duloxetine-treated group. There was also a small increase in fasting blood glucose and in total cholesterol in duloxetine-treated patients while those laboratory tests showed a slight decrease in the routine care group.

Diabetic Peripheral Neuropathic Pain

Table 2 gives the adverse reactions observed from spontaneous reporting and in placebo-controlled clinical trials (comprising a total of 6828 patients, 4199 on duloxetine and 2629 on placebo).

The most commonly observed adverse reactions in patients with diabetic neuropathic pain treated with ARICLAIM were nausea, headache, dry mouth somnolence and dizziness. However, the majority of common adverse reactions were mild to moderate, they usually started early in therapy, and most tended to subside even as therapy was continued.

Table 2: Adverse reactions

Frequency estimate: Very common ($\geq 1/10$), common ($\geq 1/100$ and <1/10), uncommon ($\geq 1/1,000$ and <1/100), rare ($\geq 1/10,000$ and <1/1,000), very rare (<1/10,000), not known (cannot be estimated from the available data).

Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Investigations				<u>.</u>	
	Weight decrease	Weight increase Creatine phosphokinase increased	Blood cholesterol increased		
Cardiac Disorder	<u>rs</u>				
	Palpitations	Tachycardia Supra- ventricular arrhythmia, mainly atrial fibrillation			
Nervous System L	Disorders				
Headache (14.3%) Somnolence (10.7%) Dizziness	Tremor Paraesthesia	Myoclonus NervousnessDis turbance in attention Lethargy	Convulsion ¹		Serotonin syndrome Extra-pyramidal symptoms Akathisia

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
(10.2%)		Dysgeusia Dyskinesia Restless legs syndrome Poor quality sleep			Psychomotor restlessness
Eye Disorders					
	Blurred vision	Mydriasis Visual disturbance	Glaucoma		
Ear and Labyrint			1		
	Tinnitus ¹	Vertigo Ear pain			
Respiratory, thore	acic and mediastine		T		
	Yawning	Throat tightness Epistaxis			
Gastrointestinal 1					
Nausea (24.3%) Dry mouth (12.8%)	Constipation Diarrhoea Vomiting Dyspepsia Flatulence	Gastroenteritis Eructation Gastritis	Stomatitis Breath odour Haematochezi-a		Gastrointestina-l haemorrhage
Renal and Urinar					
		Urinary Retention Dysuria Urinary hesitation Nocturia Polyuria Urine flow decreased	Urine odour abnormal		
Skin and Subcutar	neous Tissue Disor				
	Sweating increased Rash	Night sweats Urticaria Dermatitis contact Cold sweat Photosensitivity reactions Increased tendency to bruise			Angio-neurotic oedema Stevens- Johnson Syndrome
Musculoskeletal a	l und connective tissu	ı ve disorders			
	Musculo- skeletal pain Muscle tightness Muscle spasm	Muscle twitching	Trismus		

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Endocrine disord	ers				11110 ((12
			Hypo- thyroidism		
Metabolism and I	Nutrition Disorders		T		
	Decreased Appetite	Hyperglycemi-a (reported especially in diabetic patients)	Dehydration Hyponatremia		SIADH
Infections and inf	estations	T			
W 1 D: 1		Laryngitis			
Vascular Disorde	Flushing	Blood pressure increase Peripheral coldness Orthostatic hypotension ² Syncope ²			Hypertension Hypertensive crisis
General Disorder	l rs and Administratio	l on Site Conditions	<u> </u>		
	Fatigue Abdominal pain	Feeling abnormal Feeling cold Thirst Chills Malaise Feeling hot Gait disturbance			Chest pain
Immune system d	isorders				
·			Hyper- sensitivity disorder Anaphylactic reaction		
Hepato-biliary di	sorders				
		Elevated liver enzymes (ALT, AST, alkaline phosphatase) Hepatitis ³ Acute liver injury			Jaundice Hepatic failure
Reproductive Syst	tem and Breast Disc	orders		<u> </u>	
	Erectile dysfunction	Ejaculation disorder Ejaculation delayed Sexual dysfunction Gynaecological	Menopausal symptoms		

Very common	Common	Uncommon	Rare	Very Rare	Frequency not
					known
		haemorrhage			
Psychiatric Disor	ders				
	Insomnia	Sleep disorder	Mania		Suicidal
	Agitation	Bruxism	Hallucinations		ideation ⁵
	Libido	Disorientation	Aggression and		Suicidal ⁵
	decreased	Apathy	anger ⁴		behaviour
	Anxiety				
	Orgasm				
	abnormal				
	Abnormal				
	dreams				

Cases of convulsion and cases of tinnitus have also been reported after treatment discontinuation. ²Cases of orthostatic hypotension and syncope have been reported especially at the initiation of

³See section 4.4

Discontinuation of duloxetine (particularly when abrupt) commonly leads to withdrawal symptoms. Dizziness, sensory disturbances (including paraesthesia), sleep disturbances (including insomnia and intense dreams), agitation or anxiety, nausea and/or vomiting, tremor, headache, irritability, diarrhoea, hyperhydrosis and vertigo are the most commonly reported reactions.

Generally, for SSRIs and SNRIs, these events are mild to moderate and self-limiting, however, in some patients they may be severe and/or prolonged. It is therefore advised that when duloxetine treatment is no longer required, gradual discontinuation by dose tapering should be carried out (see sections 4.2 and 4.4).

In the 12 week acute phase of three clinical trials of duloxetine in patients with diabetic neuropathic pain, small but statistically significant increases in fasting blood glucose were observed in duloxetine-treated patients. HbA1c was stable in both duloxetine-treated and placebo-treated patients. In the extension phase of these studies, which lasted up to 52 weeks, there was an increase in HbA1c in both the duloxetine and routine care groups, but the mean increase was 0.3% greater in the duloxetine-treated group. There was also a small increase in fasting blood glucose and in total cholesterol in duloxetine-treated patients while those laboratory tests showed a slight decrease in the routine care group.

Electrocardiograms were obtained from 528 duloxetine-treated and 205 placebo-treated patients with diabetic neuropathic pain in clinical trials lasting up to 13-weeks. The heart rate-corrected QT interval in duloxetine-treated patients did not differ from that seen in placebo-treated patients. No clinically significant differences were observed for QT, PR, QRS, or QTcB measurements between duloxetine-treated and placebo-treated patients.

4.9 Overdose

Cases of overdoses, alone or in combination with other medicinal products, with duloxetine doses of 5400mg were reported. Some fatalities have occurred, primarily with mixed overdoses, but also with duloxetine alone at a dose of approximately 1000 mg. Signs and symptoms of overdose (duloxetine alone or in combination with other medicinal products) included somnolence, coma, serotonin syndrome, seizures, vomiting and tachycardia.

No specific antidote for duloxetine is known but if serotonin syndrome ensues, specific treatment (such as with cyproheptadine and/or temperature control) may be considered. A free airway should be

treatment.

⁴Cases of aggression and anger have been reported particularly early in treatment of after treatment discontinuation.

⁵Cases of suicidal ideation and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.4)

established. Monitoring of cardiac and vital signs is recommended, along with appropriate symptomatic and supportive measures. Gastric lavage may be indicated if performed soon after ingestion or in symptomatic patients. Activated charcoal may be useful in limiting absorption. Duloxetine has a large volume of distribution and forced diuresis, haemoperfusion, and exchange perfusion are unlikely to be beneficial.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other antidepressants. ATC code: N06AX21

Duloxetine is a combined serotonin (5-HT) and noradrenaline (NA) reuptake inhibitor. It weakly inhibits dopamine reuptake with no significant affinity for histaminergic, dopaminergic, cholinergic and adrenergic receptors.

In animal studies, increased levels of 5-HT and NE in the sacral spinal cord, lead to increased urethral tone via enhanced pudendal nerve stimulation to the urethral striated sphincter muscle only during the storage phase of the micturition cycle. A similar mechanism in women is believed to result in stronger urethral closure during urine storage with physical stress that could explain the efficacy of duloxetine in the treatment of women with SUI.

Duloxetine normalised pain thresholds in several preclinical models of neuropathic and inflammatory pain and attenuated pain behaviour in a model of persistent pain. The pain inhibitory action of duloxetine is believed to be a result of potentiation of descending inhibitory pain pathways within the central nervous system.

Stress Urinary Incontinence:

The efficacy of duloxetine 40 mg given twice daily in the treatment of SUI was established in four double-blind, placebo-controlled studies, that randomised 1913 women (22 to 83 years) with SUI; of these, 958 patients were randomised to duloxetine and 955 to placebo. The primary efficacy measures were Incontinence Episode Frequency (IEF) from diaries and an incontinence specific quality of life questionnaire score (I-QOL).

Incontinence Episode Frequency: in all four studies the duloxetine-treated group had a 50% or greater median decrease in IEF compared with 33% in the placebo-treated group. Differences were observed at each visit after 4 weeks (duloxetine 54%, and placebo 22%), 8 weeks (52% and 29%), and 12 weeks (52% and 33%) of medication.

In an additional study limited to patients with severe SUI, all responses with duloxetine were achieved within 2 weeks.

The efficacy of ARICLAIM has not been evaluated for longer than 3 months in placebo-controlled studies. The clinical benefit of ARICLAIM compared with placebo has not been demonstrated in women with mild SUI, defined in randomised trials as those with IEF < 14 per week. In these women, ARICLAIM may provide no benefit beyond that afforded by more conservative behavioural interventions.

Quality of Life: Incontinence Quality of Life (I-QOL) questionnaire scores were significantly improved in the duloxetine-treated patient group compared with the placebo-treated group (9.2 versus 5.9 score improvement, p<.001). Using a global improvement scale (PGI), significantly more women using duloxetine considered their symptoms of stress incontinence to be improved with treatment compared with women using placebo (64.6% versus 50.1%, p<.001).

ARICLAIM and Prior Continence Surgery: there are limited data that suggest that the benefits of ARICLAIM are not diminished in women with stress urinary incontinence who have previously undergone continence surgery.

ARICLAIM and Pelvic Floor Muscle Training (PFMT): during a 12-week blinded, randomised, controlled study, ARICLAIM demonstrated greater reductions in IEF compared with either placebo treatment or with PFMT alone. Combined therapy (duloxetine + PFMT) showed greater improvement in both pad use and condition-specific quality of life measures than ARICLAIM alone or PFMT alone.

Diabetic Peripheral Neuropathic Pain:

The efficacy of duloxetine as a treatment for diabetic neuropathic pain was established in 2 randomised, 12-week, double-blind, placebo-controlled, fixed dose studies in adults (22 to 88 years) having diabetic neuropathic pain for at least 6 months. Patients meeting diagnostic criteria for major depressive disorder were excluded from these trials. The primary outcome measure was the weekly mean of 24-hour average pain, which was collected in a daily diary by patients on an 11-point Likert scale.

In both studies, duloxetine 60 mg once daily and 60 mg twice daily significantly reduced pain compared with placebo. The effect in some patients was apparent in the first week of treatment. The difference in mean improvement between the two active treatment arms was not significant. At least 30% reported pain reduction was recorded in approximately 65% of duloxetine treated patients versus 40% for placebo. The corresponding figures for at least 50% pain reduction were 50% and 26% respectively. Clinical response rates (50% or greater improvement in pain) were analysed according to whether or not the patient experienced somnolence during treatment. For patients not experiencing somnolence, clinical response was observed in 47% of patients receiving duloxetine and 27% patients on placebo. Clinical response rates in patients experiencing somnolence were 60% on duloxetine and 30% on placebo. Patients not demonstrating a pain reduction of 30% within 60 days of treatment were unlikely to reach this level during further treatment.

In an open label long-term uncontrolled study, the pain reduction in patients responding to 8-weeks of acute treatment of ARICLAIM 60 mg once daily was maintained for a further 6-months as measured by change on the Brief Pain Inventory(BPI) 24-hour average pain item.

5.2 Pharmacokinetic properties

Duloxetine is administered as a single enantiomer. Duloxetine is extensively metabolised by oxidative enzymes (CYP1A2 and the polymorphic CYP2D6), followed by conjugation. The pharmacokinetics of duloxetine demonstrate large intersubject variability (generally 50-60%), partly due to gender, age, smoking status and CYP2D6 metaboliser status.

Duloxetine is well absorbed after oral administration with a C_{max} occurring 6 hours post dose. The absolute oral bioavailability of duloxetine ranged from 32% to 80% (mean of 50%; N=8 subjects). Food delays the time to reach the peak concentration from 6 to 10 hours and it marginally decreases the extent of absorption (approximately 11%).

Duloxetine is approximately 96% bound to human plasma proteins. Duloxetine binds to both albumin and alpha-1 acid glycoprotein. Protein binding is not affected by renal or hepatic impairment.

Duloxetine is extensively metabolised and the metabolites are excreted principally in urine. Both CYP2D6 and CYP1A2 catalyse the formation of the two major metabolites glucuronide conjugate of 4-hydroxy duloxetine and sulphate conjugate of 5-hydroxy,6-methoxy duloxetine. Based upon *in vitro* studies, the circulating metabolites of duloxetine are considered pharmacologically inactive. The pharmacokinetics of duloxetine in patients who are poor metabolisers with respect to CYP2D6 has not been specifically investigated. Limited data suggest that the plasma levels of duloxetine are higher in these patients.

The elimination half-life of duloxetine after an oral dose ranges from 8 to 17 hours (mean of 12 hours). After an intravenous dose the plasma clearance of duloxetine ranges from 22 l/hr to 46 l/hr (mean of 36 l/hr) After an oral dose the apparent plasma clearance of duloxetine ranges from 33 to 261 l/hr (mean 101 l/hr).

Special populations:

Gender: pharmacokinetic differences have been identified between males and females (apparent plasma clearance is approximatively 50% lower in females). Based upon the overlap in the range of clearance, gender-based pharmacokinetic differences do not justify the recommendation for using a lower dose for female patients.

Age: pharmacokinetic differences have been identified between younger and elderly females (\geq 65 years) (AUC increases by about 25% and half-life is about 25% longer in the elderly), although the magnitude of these changes is not sufficient to justify adjustments to the dose. As a general recommendation, caution should be exercised when treating the elderly (see sections 4.2 and 4.4).

Renal impairment: end stage renal disease (ESRD) patients receiving dialysis had a 2-fold higher duloxetine C_{max} and AUC values compared to healthy subjects. Pharmacokinetic data on duloxetine is limited in patients with mild or moderate renal impairment.

Hepatic insufficiency: moderate liver disease (Child Pugh Class B) affected the pharmacokinetics of duloxetine. Compared with healthy subjects, the apparent plasma clearance of duloxetine was 79% lower, the apparent terminal half-life was 2.3 times longer, and the AUC was 3.7 times higher in patients with moderate liver disease. The pharmacokinetics of duloxetine and its metabolites have not been studied in patients with mild or severe hepatic insufficiency.

Nursing mothers: The disposition of duloxetine was studied in 6 lactating women who were at least 12-weeks postpartum. Duloxetine is detected in breast milk, and steady-state concentrations in breast milk are about one-fourth those in plasma. The amount of duloxetine in breast milk is approximately 7 μg/day while on 40 mg twice daily dosing. Lactation did not influence duloxetine pharmacokinetics.

5.3 Preclinical safety data

Duloxetine was not genotoxic in a standard battery of tests and was not carcinogenic in rats. Multinucleated cells were seen in the liver in the absence of other histopathological changes in the rat carcinogenicity study. The underlying mechanism and the clinical relevance are unknown.

Female mice receiving duloxetine for 2 years had an increased incidence of hepatocellular adenomas and carcinomas at the high dose only (144 mg/kg/day), but these were considered to be secondary to hepatic microsomal enzyme induction. The relevance of this mouse data to humans is unknown. Female rats receiving duloxetine before and during mating and early pregnancy had a decrease in maternal food consumption and body weight, oestrous cycle disruption, decreased live birth indices and progeny survival, and progeny growth retardation at systemic exposure levels estimated to be at the most at maximum clinical exposure (AUC). In an embryotoxicity study in the rabbit, a higher incidence of cardiovascular and skeletal malformations was observed at systemic exposure levels below the maximum clinical exposure (AUC). No malformations were observed in another study testing a higher dose of a different salt of duloxetine. In pre/postnatal toxicity study in the rat, duloxetine induced adverse behavioural effects in the offspring at systemic exposures levels below maximum clinical exposure (AUC).

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Capsule content:

Hypromellose.

Hypromellose acetate succinate Sucrose Sugar spheres Talc Titanium dioxide (E171) Triethyl citrate.

Capsule shell:

Gelatin Sodium Lauryl Sulfate Titanium Dioxide (E171) Indigo Carmine (E132) Edible Black Ink.

Edible Ink: Black Iron Oxide-Synthetic (E172) Propylene glycol Shellac

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years.

6.4 Special precautions for storage

Store in the original package in order to protect from moisture. Do not store above 30°C.

6.5 Nature and contents of container

Polyvinylchloride (PVC), Polyethylene (PE), and Polychlorotrifluoroethylene (PCTFE) blister sealed with an aluminium foil.

Packs of 28 and 56 capsules.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

Eli Lilly Nederland BV, Grootslag 1-5, NL-3991 RA Houten, The Netherlands.

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/04/283/001 EU/1/04/283/007

9.	DATE OF FIRST A	UTHORISATION/RENEWAL OF	F THE AUTHORISATION
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11 August 2004

10. DATE OF REVISION OF THE TEXT

1. NAME OF THE MEDICINAL PRODUCT

ARICLAIM 30 mg hard gastro-resistant capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each capsule contains 30 mg of duloxetine (as hydrochloride)

Excipients: sucrose 8.6 mg.

For a full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard gastro-resistant capsule.

Opaque white body, imprinted with '30 mg' and an opaque blue cap, imprinted with '9543'.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Treatment of diabetic peripheral neuropathic pain in adults.

4.2 Posology and method of administration

For oral use.

Adults

The starting and recommended maintenance dose is 60 mg daily with or without food. Dosages above 60 mg once daily, up to a maximum dose of 120 mg per day administered in evenly divided doses, have been evaluated from a safety perspective in clinical trials. The plasma concentration of duloxetine displays large inter-individual variability (see 5.2). Hence, some patients that respond insufficiently to 60 mg may benefit from a higher dose.

Response to treatment should be evaluated after 2 months. In patients with inadequate initial response, additional response after this time is unlikely.

The therapeutic benefit should be reassessed regularly (at least every three months) (see Section 5.1).

Elderly

No dosage adjustment is recommended for elderly patients solely on the basis of age. However, caution should be exercised when treating the elderly (see section 5.2).

Children and adolescents

There is no experience in children (see section 4.4).

Hepatic impairment

ARICLAIM should not be used in patients with liver disease resulting in hepatic impairment (see sections 4.3 and 5.2).

Renal insufficiency

No dosage adjustment is necessary for patients with mild or moderate renal dysfunction (creatinine clearance 30 to 80 ml/min). ARICLAIM should not be used in patients with severe renal impairment (creatinine clearance <30 ml/min; see section 4.3)..

Discontinuation of treatment

Abrupt discontinuation should be avoided. When stopping treatment with ARICLAIM the dose should be gradually reduced over a period of at least one to two weeks in order to reduce the risk of withdrawal reactions (see sections 4.4 and 4.8). If intolerable symptoms occur following a decrease in the dose or upon discontinuation of treatment, then resuming the previously prescribed dose may be considered. Subsequently, the physician may continue decreasing the dose, but at a more gradual rate.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients.

Concomitant use of ARICLAIM with nonselective, irreversible Monoamine Oxidase Inhibitors (MAOIs) is contraindicated (see section 4.5).

Liver disease resulting in hepatic impairment (see section 5.2).

ARICLAIM should not be used in combination with fluvoxamine, ciprofloxacin or enoxacine (i.e. potent CYP1A2 inhibitors) since the combination results in elevated plasma concentrations of duloxetine (see section 4.5).

Severe renal impairment (creatinine clearance <30 ml/min) (see section 4.4).

The initiation of treatment with ARICLAIM is contraindicated in patients with uncontrolled hypertension that could expose patients to a potential risk of hypertensive crisis (see sections 4.4 and 4.8).

4.4 Special warnings and precautions for use

Mania and seizures

ARICLAIM should be used with caution in patients with a history of mania or a diagnosis of bipolar disorder, and/or seizures.

Mydriasis

Mydriasis has been reported in association with duloxetine, therefore, caution should be used when prescribing ARICLAIM to patients with increased intraocular pressure, or those at risk of acute narrow-angle glaucoma.

Blood pressure and heart rate

Duloxetine has been associated with an increase in blood pressure and clinically significant hypertension in some patients. This may be due to the noradrenergic effect of duloxetine. Cases of hypertensive crisis have been reported with duloxetine, especially in patients with pre-existing hypertension. Therefore, in patients with known hypertension and/or other cardiac disease, blood pressure monitoring is recommended, especially during the first month of treatment. Duloxetine should be used with caution in patients whose conditions could be compromised by an increased heart rate or by an increase in blood pressure. Caution should also be exercised when duloxetine is used with medicinal products that may impair its metabolism (see section 4.5). For patients who experience a sustained increase in blood pressure while receiving duloxetine either dose reduction or gradual discontinuation should be considered (see section 4.8). In patients with uncontrolled hypertension duloxetine should not be initiated (see section 4.3).

Renal impairment

Increased plasma concentrations of duloxetine occur in patients with severe renal impairment on haemodialysis (creatinine clearance <30 ml/min). For patients with severe renal impairment, see section 4.3. ee section 4.2 for information on patients with mild or moderate renal dysfunction.

Use with antidepressants

Caution should be exercised when using ARICLAIM in combination with antidepressants. In particular the combination with selective reversible MAOIs is not recommended.

St John's wort

Undesirable effects may be more common during concomitant use of ARICLAIM and herbal preparations containing St John's wort (Hypericum perforatum).

Depression, suicidal ideation and behaviour

Although ARICLAIM is not indicated for the treatment of depression, its active ingredient (duloxetine) also exists as an antidepressant medication. Depression is associated with an increased risk of suicidal thoughts, self harm and suicide (suicide-related events). This risk persists until significant remission occurs. As improvement may not occur during the first few weeks or more of treatment, patients should be closely monitored until such improvement occurs. It is general clinical experience that the risk of suicide may increase in the early stages of recovery. Patients with a history of suicide-related events or those exhibiting a significant degree of suicidal thoughts prior to commencement of treatment are known to be at a greater risk of suicidal thoughts or suicidal behaviour, and should receive careful monitoring during treatment. A meta-analysis of placebo-controlled clinical trials of antidepressant medicinal products in psychiatric disorders showed an increased risk of suicidal behaviour with antidepressants compared to placebo in patients less than 25 years old.

Cases of suicidal thoughts and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.8). Physicians should encourage patients to report any distressing thoughts or feelings or depressive symptoms at any time. If while on ARICLAIM therapy, the patient develops agitation or depressive symptoms, specialised medical advice should be sought, as depression is a serious medical condition. If a decision to initiate antidepressant pharmacological therapy is taken, the gradual discontinuation of ARICLAIM is recommended (see section 4.2).

Use in children and adolescents under 18 years of age

No clinical trials have been conducted with duloxetine in paediatric populations. ARICLAIM should not be used in the treatment of children and adolescents under the age of 18 years. Suicide-related behaviours (suicide attempts and suicidal thoughts), and hostility (predominantly aggression, oppositional behaviour and anger), were more frequently observed in clinical trials among children and adolescents treated with antidepressants compared to those treated with placebo. If, based on clinical need, a decision to treat is nevertheless taken, the patient should be carefully monitored for the appearance of suicidal symptoms. In addition, long-term safety data in children and adolescents concerning growth, maturation and cognitive and behavioural development are lacking.

Haemorrhage

There have been reports of bleeding abnormalities, such as ecchymoses, purpura and gastrointestinal haemorrhage with selective serotonin reuptake inhibitors (SSRIs) and serotonin/noradrenaline reuptake inhibitors (SNRIs). Caution is advised in patients taking anticoagulants and/or medicinal products known to affect platelet function, and in patients with known bleeding tendencies.

Hyponatremia

Hyponatremia has been reported rarely, predominantly in the elderly, when administering ARICLAIM. Caution is required in patients at increased risk for hyponatremia; such as elderly, cirrhotic, or dehydrated patients or patients treated with diuretics. Hyponatremia may be due to a syndrome of inappropriate anti-diuretic hormone secretion (SIADH).

Discontinuation of treatment

Withdrawal symptoms when treatment is discontinued are common, particularly if discontinuation is abrupt (see section 4.8). In clinical trials adverse events seen on abrupt treatment discontinuation occurred in approximately 45% of patients treated with ARICLAIM and 23% of patients taking placebo.

The risk of withdrawal symptoms seen with SSRI's and SNRI's may be dependent on several factors including the duration and dose of therapy and the rate of dose reduction. The most commonly reported reactions are listed in section 4.8. Generally these symptoms are mild to moderate, however, in some patients they may be severe in intensity. They usually occur within the first few days of discontinuing treatment, but there have been very rare reports of such symptoms in patients who have inadvertently missed a dose. Generally these symptoms are self-limiting and usually resolve within 2 weeks, though in some individuals they may be prolonged (2-3 months or more). It is therefore advised that duloxetine should be gradually tapered when discontinuing treatment over a period of no less than 2 weeks, according to the patient's needs (see section 4.2).

Akathisia/psychomotor restlessness

The use of duloxetine has been associated with the development of akathisia, characterised by a subjectively unpleasant or distressing restlessness and need to move often accompanied by an inability to sit or stand still. This is most likely to occur within the first few weeks of treatment. In patients who develop these symptoms, increasing the dose may be detrimental.

Medicinal products containing duloxetine

Duloxetine is used under different trademarks in several indications (treatment of diabetic neuropathic pain, major depressive episodes as well as stress urinary incontinence). The use of more than one of these products concomitantly should be avoided.

Hepatitis/increased liver enzymes

Cases of liver injury, including severe elevations of liver enzymes (>10 times upper limit of normal), hepatitis and jaundice have been reported with duloxetine (see section 4.8). Most of them occurred during the first months of treatment. The pattern of liver damage was predominantly hepatocellular. Duloxetine should be used with caution in patients treated with other medicinal products associated with hepatic injury.

Sucrose

ARICLAIM gastro-resistant capsules contain sucrose. Patients with rare hereditary problems of fructose intolerance, glucose-galactose malabsorption or sucrose-isomaltase insufficiency should not take this medicine.

4.5 Interaction with other medicinal products and other forms of interaction

CNS medicinal products: the risk of using duloxetine in combination with other CNS-active medicinal products has not been systematically evaluated, except in the cases described in this section. Consequently, caution is advised when ARICLAIM is taken in combination with other centrally acting medicinal products and substances including alcohol and sedative medicinal products (e.g. benzodiazepines, morphinomimetics, antipsychotics, phenobarbital, sedative antihistamines).

Monoamine Oxidase Inhibitors (MAOIs): due to the risk of serotonin syndrome, ARICLAIM should not be used in combination with nonselective irreversible monoamine oxidase inhibitors (MAOIs), or within at least 14 days of discontinuing treatment with an MAOI. Based on the half-life of duloxetine, at least 5 days should be allowed after stopping ARICLAIM before starting an MAOI (see section 4.3).

For selective, reversible MAOIs, like moclobemide, the risk of serotonin syndrome is lower. However, the concomitant use of ARICLAIM with selective, reversible MAOIs is not recommended (see section 4.4).

Serotonin syndrome: in rare cases, serotonin syndrome has been reported in patients using SSRIs (e.g. paroxetine, fluoxetine) concomitantly with serotonergic medicinal products. Caution is advisable if ARICLAIM is used concomitantly with serotonergic antidepressants like SSRIs, tricyclics like clomipramine or amitriptyline, St John's wort (Hypericum perforatum), venlafaxine or triptans, tramadol, pethidine and tryptophan.

Effect of duloxetine on other medicinal products

Medicinal products metabolised by CYP1A2: The pharmacokinetics of theophylline, a CYP1A2 substrate, were not significantly affected by co-administration with duloxetine (60 mg twice daily). Medicinal products metabolised by CYP2D6: Duloxetine is a moderate inhibitor of CYP2D6. When duloxetine was administered at a dose of 60 mg twice daily with a single dose of desipramine, a CYP2D6 substrate, the AUC of desipramine increased 3-fold. The co-administration of duloxetine (40 mg twice daily) increases steady state AUC of tolterodine (2 mg twice daily) by 71 %, but does not affect the pharmacokinetics of its active 5-hydroxyl metabolite and no dosage adjustment is recommended. Caution is advised if ARICLAIM is co-administered with medicinal products that are predominantly metabolised by CYP2D6 (risperidone, tricyclic antidepressants [TCAs] such as nortriptyline, amitriptyline, and imipramine) particularly if they have a narrow therapeutic index (such as flecainide, propafenone and metoprolol).

Oral contraceptives and other steroidal agents: results of *in vitro* studies demonstrate that duloxetine does not induce the catalytic activity of CYP3A. Specific *in vivo* drug interaction studies have not been performed.

Anticoagulants and antiplatelet agents: Caution should be exercised when duloxetine is combined with oral anticoagulants or antiplatelet agents due to a potential increased risk of bleeding. Furthermore, increases in INR values have been reported when duloxetine was co-administered with warfarin.

Effects of other medicinal products on duloxetine

Antacids and H2 antagonists: co-administration of duloxetine with aluminium- and magnesium-containing antacids or duloxetine with famotidine had no significant effect on the rate or extent of duloxetine absorption after administration of a 40 mg oral dose.

Inhibitors of CYP 1A2: because CYP1A2 is involved in duloxetine metabolism, concomitant use of duloxetine with potent inhibitors of CYP1A2 is likely to result in higher concentrations of duloxetine. Fluvoxamine (100 mg once daily), a potent inhibitor of CYP1A2, decreased the apparent plasma clearance of duloxetine by about 77% and increased AUC_{0-t} 6-fold. Therefore ARICLAIM should not be administered in combination with potent inhibitors of CYP1A2 like fluvoxamine (see section 4.3).

Inducers of CYP1A2: Population pharmacokinetic analyses have shown that smokers have almost 50% lower plasma concentrations of duloxetine compared with non-smokers.

4.6 Pregnancy and lactation

Pregnancy

There are no data on the use of duloxetine in pregnant women. Studies in animals have shown reproductive toxicity at systemic exposure levels (AUC) of duloxetine lower than the maximum clinical exposure (see section 5.3).

The potential risk for humans is unknown. As with other serotonergic medicinal products, discontinuation symptoms may occur in the neonate after maternal duloxetine use near term. ARICLAIM should be used in pregnancy only if the potential benefit justifies the potential risk to the foetus. Women should be advised to notify their physician if they become pregnant, or intend to become pregnant, during therapy.

Breast feeding

Duloxetine is very weakly excreted into human milk based on a study of 6 lactating patients, who did not breast feed their children . The estimated daily infant dose on a mg/kg basis is approximately 0.14% of the maternal dose (see section 5.2). As the safety of duloxetine in infants is not known, the use of ARICLAIM while breast-feeding is not recommended.

4.7 Effects on ability to drive and use machines

No studies on the effects on the ability to drive and use machines have been performed. ARICLAIM may be associated with sedation and dizziness. Patients should be instructed that if they experience sedation or dizziness they should avoid potentially hazardous tasks such as driving or operating machinery.

4.8 Undesirable effects

Table 1 gives the adverse reactions observed from spontaneous reporting and in placebo-controlled clinical trials (comprising a total of 6828 patients, 4199 on duloxetine and 2629 on placebo). The most commonly observed adverse reactions in patients with diabetic neuropathic pain treated with ARICLAIM were: nausea; headache, dry mouth and somnolence and dizziness.

Table 1: Adverse reactions

Frequency estimate: Very common ($\geq 1/10$), common ($\geq 1/100$ and <1/10), uncommon ($\geq 1/1,000$ and <1/100), rare ($\geq 1/10,000$ and <1/1,000), very rare (<1/10,000), not known (cannot be estimated from the available data).

Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Investigations				•	•
Cardiac Disorde	Weight decrease	Weight increase Creatine phosphokinase increased	Blood cholesterol increased		
	Palpitations	Tachycardia Supra- ventricular arrhythmia, mainly atrial fibrillation			
Nervous System I)	Γ	T	
Headache (14.3%) Somnolence (10.7%) Dizziness (10.2%)	Tremor Paraesthesia	Myoclonus NervousnessDis turbance in attention Lethargy Dysgeusia Dyskinesia Restless legs syndrome Poor quality sleep	Convulsion ¹		Serotonin syndrome Extra-pyramidal symptoms Akathisia Psychomotor restlessness
Eye Disorders				l .	
	Blurred vision	Mydriasis Visual disturbance	Glaucoma		
Ear and Labyrint	h Disorders			1	
	Tinnitus ¹	Vertigo Ear pain			

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Respiratory, thor	acic and mediastin	al disorders			
	Yawning	Throat tightness Epistaxis			
Gastrointestinal I					
Nausea (24.3%)	Constipation	Gastroenteritis	Stomatitis		Gastrointestinal
Dry mouth	Diarrhoea	Eructation	Breath odour		haemorrhage
(12.8%)	Vomiting	Gastritis	Haematochezia		
	Dyspepsia				
D1 1 IIi	Flatulence				
Renal and Urina	ry Disoraers	Lininger	Urine odour		<u> </u>
		Urinary Retention	abnormal		
			abnormai		
		Dysuria Urinary			
		hesitation			
l		Nocturia			
		Polyuria			
		Urine flow			
		decreased			
Skin and Subcuta	neous Tissue Disor		<u>ı</u>		_ 1
21111 01110 2110 01110	Sweating	Night sweats			Angio-neurotic
	increased	Urticaria			oedema
	Rash	Dermatitis			Stevens-
		contact			Johnson
		Cold sweat			Syndrome
		Photo-			
		sensitivity			
		reactions			
		Increased			
		tendency to			
		bruise			
Musculoskeletal d	and connective tissi	ue disorders			
1	Musculo-	Muscle	Trismus		
	skeletal pain	twitching			
	Muscle				
	tightness				
	Muscle spasm				
Endocrine disord	lers	1	111		
			Hypo-		
M - 4 = 1 - 1: 1:	Nadariti an Dia and an		thyroidism		
meiavousm and l	Nutrition Disorders Decreased		Dehydration		SIADH
	Appetite	Hyperglycemi-a (reported	Dehydration Hyponatremia		ылип
	Appenie	especially in	Пуропансина		
		diabetic			
		patients)			
Infections and in	festations	1 Parietiu)	1		_1
.,		Laryngitis			
Vascular Disorde	ers	, <i>, , ,</i>	<u>ı</u> l		_1
	Flushing	Blood pressure			Hypertension
		increase			Hypertensive

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
		coldness Orthostatic hypotension ² Syncope ²			
General Disorder	rs and Administration	on Site Conditions			
	Fatigue Abdominal pain	Feeling abnormal Feeling cold Thirst Chills Malaise Feeling hot Gait disturbance			Chest pain
Immune system di	isorders				
			Hyper- sensitivity disorder Anaphylactic reaction		
Hepato-biliary di	sorders	T1 (11)	Г		T 1.
		Elevated liver enzymes (ALT, AST, alkaline phosphatase) Hepatitis ³ Acute liver injury			Jaundice Hepatic failure
Reproductive Syst	tem and Breast Disc	orders			
	Erectile dysfunction	Ejaculation disorder Ejaculation delayed Sexual dysfunction Gynaecological haemorrhage	Menopausal symptoms		
Psychiatric Disor		G1 1: 1			
	Insomnia Agitation Libido decreased Anxiety Orgasm abnormal Abnormal dreams	Sleep disorder Bruxism Disorientation Apathy	Mania Hallucinations Aggression and anger ⁴		Suicidal ideation ⁵ Suicidal ⁵ behaviour

Cases of convulsion and cases of tinnitus have also been reported after treatment discontinuation.

Cases of orthostatic hypotension and syncope have been reported especially at the initiation of treatment.

Discontinuation of duloxetine (particularly when abrupt) commonly leads to withdrawal symptoms. Dizziness, sensory disturbances (including paraesthesia), sleep disturbances (including insomnia and intense dreams), agitation or anxiety, nausea and/or vomiting, tremor ,headache, irritability, diarrhoea, hyperhydrosis and vertigo are the most commonly reported reactions.

Generally, for SSRIs and SNRIs, these events are mild to moderate and self-limiting, however, in some patients they may be severe and/or prolonged. It is therefore advised that when duloxetine treatment is no longer required, gradual discontinuation by dose tapering should be carried out (see sections 4.2 and 4.4).

In the 12 week acute phase of three clinical trials of duloxetine in patients with diabetic neuropathic pain, small but statistically significant increases in fasting blood glucose were observed in duloxetine-treated patients. HbA1c was stable in both duloxetine-treated and placebo-treated patients. In the extension phase of these studies, which lasted up to 52 weeks, there was an increase in HbA1c in both the duloxetine and routine care groups, but the mean increase was 0.3% greater in the duloxetine-treated group. There was also a small increase in fasting blood glucose and in total cholesterol in duloxetine-treated patients while those laboratory tests showed a slight decrease in the routine care group.

Electrocardiograms were obtained from 528 duloxetine-treated and 205 placebo-treated patients with diabetic neuropathic pain in clinical trials lasting up to 13-weeks. The heart rate-corrected QT interval in duloxetine-treated patients did not differ from that seen in placebo-treated patients. No clinically significant differences were observed for QT, PR, QRS, or QTcB measurements between duloxetine-treated and placebo-treated patients.

4.9 Overdose

Cases of overdoses, alone or in combination with other medicinal products, with duloxetine doses of 5400mg were reported. Some fatalities have occurred, primarily with mixed overdoses, but also with duloxetine alone at a dose of approximately 1000 mg. Signs and symptoms of overdose (duloxetine alone or in combination with other medicinal products) included somnolence, coma, serotonin syndrome, seizures, vomiting and tachycardia.

No specific antidote is known for duloxetine but if serotonin syndrome ensues, specific treatment (such as with cyproheptadine and/or temperature control) may be considered. A free airway should be established. Monitoring of cardiac and vital signs is recommended, along with appropriate symptomatic and supportive measures. Gastric lavage may be indicated if performed soon after ingestion or in symptomatic patients. Activated charcoal may be useful in limiting absorption. Duloxetine has a large volume of distribution and forced diuresis, haemoperfusion, and exchange perfusion are unlikely to be beneficial.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other antidepressants. ATC code: N06AX21.

Duloxetine is a combined serotonin (5-HT) and noradrenaline (NA) reuptake inhibitor. It weakly inhibits dopamine reuptake with no significant affinity for histaminergic, dopaminergic, cholinergic and adrenergic receptors. Duloxetine dose-dependently increases extracellular levels of serotonin and noradrenaline in various brain areas of animals.

³See section 4.4

⁴Cases of aggression and anger have been reported particularly early in treatment of after treatment discontinuation.

⁵Cases of suicidal ideation and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.4)

Duloxetine normalised pain thresholds in several preclinical models of neuropathic and inflammatory pain and attenuated pain behaviour in a model of persistent pain. The pain inhibitory action of duloxetine is believed to be a result of potentiation of descending inhibitory pain pathways within the central nervous system.

The efficacy of duloxetine as a treatment for diabetic neuropathic pain was established in 2 randomised, 12-week, double-blind, placebo-controlled, fixed dose studies in adults (22 to 88 years) having diabetic neuropathic pain for at least 6 months. Patients meeting diagnostic criteria for major depressive disorder were excluded from these trials. The primary outcome measure was the weekly mean of 24-hour average pain, which was collected in a daily diary by patients on an 11-point Likert scale.

In both studies, duloxetine 60 mg once daily and 60 mg twice daily significantly reduced pain compared with placebo. The effect in some patients was apparent in the first week of treatment. The difference in mean improvement between the two active treatment arms was not significant. At least 30% reported pain reduction was recorded in approximately 65% of duloxetine treated patients versus 40% for placebo. The corresponding figures for at least 50% pain reduction were 50% and 26% respectively. Clinical response rates (50% or greater improvement in pain) were analysed according to whether or not the patient experienced somnolence during treatment. For patients not experiencing somnolence, clinical response was observed in 47% of patients receiving duloxetine and 27% patients on placebo. Clinical response rates in patients experiencing somnolence were 60% on duloxetine and 30% on placebo. Patients not demonstrating a pain reduction of 30% within 60 days of treatment were unlikely to reach this level during further treatment.

In an open label long-term uncontrolled study, the pain reduction in patients responding to 8-weeks of acute treatment of ARICLAIM 60 mg once daily was maintained for a further 6-months as measured by change on the Brief Pain Inventory(BPI) 24-hour average pain item.

5.2 Pharmacokinetic properties

Duloxetine is administered as a single enantiomer. Duloxetine is extensively metabolised by oxidative enzymes (CYP1A2 and the polymorphic CYP2D6), followed by conjugation. The pharmacokinetics of duloxetine demonstrate large intersubject variability (generally 50-60%), partly due to gender, age, smoking status and CYP2D6 metaboliser status.

Duloxetine is well absorbed after oral administration with a Cmax occurring 6 hours post dose. The absolute oral bioavailability of duloxetine ranged from 32% to 80% (mean of 50%). Food delays the time to reach the peak concentration from 6 to 10 hours and it marginally decreases the extent of absorption (approximately 11%). These changes do not have any clinical significance. Duloxetine is approximately 96% bound to human plasma proteins. Duloxetine binds to both albumin and alphalacid glycoprotein. Protein binding is not affected by renal or hepatic impairment.

Duloxetine is extensively metabolised and the metabolites are excreted principally in urine. Both cytochromes P450-2D6 and 1A2 catalyse the formation of the two major metabolites glucuronide conjugate of 4-hydroxy duloxetine and sulphate conjugate of 5hydroxy,6-methoxy duloxetine. Based upon *in vitro* studies, the circulating metabolites of duloxetine are considered pharmacologically inactive. The pharmacokinetics of duloxetine in patients who are poor metabolisers with respect to CYP2D6 has not been specifically investigated. Limited data suggest that the plasma levels of duloxetine are higher in these patients.

The elimination half-life of duloxetine ranges from 8 to 17 hours (mean of 12 hours). After an intravenous dose the plasma clearance of duloxetine ranges from 22 l/hr to 46 l/hr (mean of 36 l/hr). After an oral dose the apparent plasma clearance of duloxetine ranges from 33 to 261 l/hr (mean 101 l/hr).

Special populations:

Gender: pharmacokinetic differences have been identified between males and females (apparent plasma clearance is approximatively 50% lower in females). Based upon the overlap in the range of clearance, gender-based pharmacokinetic differences do not justify the recommendation for using a lower dose for female patients.

Age: pharmacokinetic differences have been identified between younger and elderly females (\geq 65 years) (AUC increases by about 25% and half-life is about 25% longer in the elderly), although the magnitude of these changes is not sufficient to justify adjustments to the dose. As a general recommendation, caution should be exercised when treating the elderly (see sections 4.2 and 4.4).

Renal impairment: end stage renal disease (ESRD) patients receiving dialysis had 2-fold higher duloxetine Cmax and AUC values compared with healthy subjects. Pharmacokinetic data on duloxetine is limited in patients with mild or moderate renal impairment.

Hepatic impairment: moderate liver disease (Child Pugh Class B) affected the pharmacokinetics of duloxetine. Compared with healthy subjects, the apparent plasma clearance of duloxetine was 79% lower, the apparent terminal half-life was 2.3 times longer, and the AUC was 3.7 times higher in patients with moderate liver disease. The pharmacokinetics of duloxetine and its metabolites have not been studied in patients with mild or severe hepatic insufficiency.

Nursing mothers: The disposition of duloxetine was studied in 6 lactating women who were at least 12-weeks postpartum. Duloxetine is detected in breast milk, and steady-state concentrations in breast milk are about one-fourth those in plasma. The amount of duloxetine in breast milk is approximately 7 μg/day while on 40 mg twice daily dosing. Lactation did not influence duloxetine pharmacokinetics.

5.3 Preclinical safety data

Duloxetine was not genotoxic in a standard battery of tests and was not carcinogenic in rats. Multinucleated cells were seen in the liver in the absence of other histopathological changes in the rat carcinogenicity study. The underlying mechanism and the clinical relevance are unknown. Female mice receiving duloxetine for 2 years had an increased incidence of hepatocellular adenomas and carcinomas at the high dose only (144 mg/kg/day), but these were considered to be secondary to hepatic microsomal enzyme induction. The relevance of this mouse data to humans is unknown. Female rats receiving duloxetine (45 mg/kg/day) before and during mating and early pregnancy had a decrease in maternal food consumption and body weight, oestrous cycle disruption, decreased live birth indices and progeny survival, and progeny growth retardation at systemic exposure levels estimated to be at the most at maximum clinical exposure (AUC). In an embryotoxicity study in the rabbit, a higher incidence of cardiovascular and skeletal malformations was observed at systemic exposure levels below the maximum clinical exposure (AUC). No malformations were observed in another study testing a higher dose of a different salt of duloxetine. In prenatal/postnatal toxicity studies in the rat, duloxetine induced adverse behavioural effects in the offspring at exposures below maximum clinical exposure (AUC).

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Capsule content:

Hypromellose.
Hypromellose acetate succinate
Sucrose
Sugar spheres
Talc
Titanium dioxide (E171)
Triethyl citrate.

Capsule shell:

30 mg: Gelatin Sodium Lauryl Sulfate Titanium Dioxide (E171) Indigo Carmine (E132) Edible Green Ink

Edible Green Ink contains: Black Iron Oxide-Synthetic (E172) Yellow Iron Oxide-Synthetic (E172) Propylene glycol Shellac.

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years.

6.4 Special precautions for storage

Store in the original package in order to protect from moisture. Do not store above 30 C.

6.5 Nature and contents of container

Polyvinylchloride (PVC), Polyethylene (PE), and Polychlorotrifluoroethylene (PCTFE) blister sealed with an aluminium foil.

ARICLAIM 30 mg is available in packs of 7, 28 and 98 capsules.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

Eli Lilly Nederland BV, Grootslag 1-5, NL-3991 RA Houten, The Netherlands.

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/04/283/008 EU/1/04/283/009 EU/1/04/283/010

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

10. DATE OF REVISION OF THE TEXT

1. NAME OF THE MEDICINAL PRODUCT

ARICLAIM 60 mg hard gastro-resistant capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each capsule contains 60 mg of duloxetine (as hydrochloride).

Excipients: sucrose 17.2 mg.

For a full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard gastro-resistant capsule.

Opaque green body, imprinted with '60 mg' and an opaque blue cap, imprinted with '9542'.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Treatment of diabetic peripheral neuropathic pain in adults.

4.2 Posology and method of administration

For oral use.

Adults

The starting and recommended maintenance dose is 60 mg daily with or without food. Dosages above 60 mg once daily, up to a maximum dose of 120 mg per day administered in evenly divided doses, have been evaluated from a safety perspective in clinical trials. The plasma concentration of duloxetine displays large inter-individual variability (see 5.2). Hence, some patients that respond insufficiently to 60 mg may benefit from a higher dose.

Response to treatment should be evaluated after 2 months. In patients with inadequate initial response, additional response after this time is unlikely.

The therapeutic benefit should be reassessed regularly (at least every three months) (see Section 5.1).

Elderly

Diabetic Peripheral Neuropathic Pain: No dosage adjustment is recommended for elderly patients solely on the basis of age. However, caution should be exercised when treating the elderly (see section 5.2).

Children and adolescents

There is no experience in children (see section 4.4).

Hepatic impairment

ARICLAIM should not be used in patients with liver disease resulting in hepatic impairment (see sections 4.3 and 5.2).

Renal insufficiency

No dosage adjustment is necessary for patients with mild or moderate renal dysfunction (creatinine clearance 30 to 80 ml/min). ARICLAIM should not be used in patients with severe renal impairment (creatinine clearance <30 ml/min; see section 4.3).

Discontinuation of treatment

Abrupt discontinuation should be avoided. When stopping treatment with ARICLAIM the dose should be gradually reduced over a period of at least one to two weeks in order to reduce the risk of withdrawal reactions (see sections 4.4 and 4.8). If intolerable symptoms occur following a decrease in the dose or upon discontinuation of treatment, then resuming the previously prescribed dose may be considered. Subsequently, the physician may continue decreasing the dose, but at a more gradual rate.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients.

Concomitant use of ARICLAIM with nonselective, irreversible Monoamine Oxidase Inhibitors (MAOIs) is contraindicated (see section 4.5).

Liver disease resulting in hepatic impairment (see section 5.2).

ARICLAIM should not be used in combination with fluvoxamine, ciprofloxacin or enoxacine (i.e. potent CYP1A2 inhibitors) since the combination results in elevated plasma concentrations of duloxetine (see section 4.5).

Severe renal impairment (creatinine clearance <30 ml/min) (see section 4.4).

The initiation of treatment with ARICLAIM is contraindicated in patients with uncontrolled hypertension that could expose patients to a potential risk of hypertensive crisis (see sections 4.4 and 4.8).

4.4 Special warnings and precautions for use

Mania and seizures

ARICLAIM should be used with caution in patients with a history of mania or a diagnosis of bipolar disorder, and/or seizures.

Mydriasis

Mydriasis has been reported in association with duloxetine, therefore, caution should be used when prescribing ARICLAIM to patients with increased intraocular pressure, or those at risk of acute narrow-angle glaucoma.

Blood pressure and heart rate

Duloxetine has been associated with an increase in blood pressure and clinically significant hypertension in some patients. This may be due to the noradrenergic effect of duloxetine. Cases of hypertensive crisis have been reported with duloxetine, especially in patients with pre-existing hypertension. Therefore,in patients with known hypertension and/or other cardiac disease, blood pressure monitoring is recommended, especially during the first month of treatment. Duloxetine should be used with caution in patients whose conditions could be compromised by an increased heart rate or by an increase in blood pressure. Caution should also be exercised when duloxetine is used with medicinal products that may impair its metabolism (see section 4.5). For patients who experience a sustained increase in blood pressure while receiving duloxetine either dose reduction or gradual discontinuation should be considered (see section 4.8). In patients with uncontrolled hypertension duloxetine should not be initiated (see section 4.3).

Renal impairment

Increased plasma concentrations of duloxetine occur in patients with severe renal impairment on haemodialysis (creatinine clearance <30 ml/min). For patients with severe renal impairment, see section 4.3. See section 4.2 for information on patients with mild or moderate renal dysfunction.

Use with antidepressants

Caution should be exercised when using ARICLAIM in combination with antidepressants. In particular the combination with selective reversible MAOIs is not recommended.

St John's wort

Undesirable effects may be more common during concomitant use of ARICLAIM and herbal preparations containing St John's wort (Hypericum perforatum).

Depression, suicidal ideation and behaviour

Although ARICLAIM is not indicated for the treatment of depression, its active ingredient (duloxetine) also exists as an antidepressant medication. Depression is associated with an increased risk of suicidal thoughts, self harm and suicide (suicide-related events). This risk persists until significant remission occurs. As improvement may not occur during the first few weeks or more of treatment, patients should be closely monitored until such improvement occurs. It is general clinical experience that the risk of suicide may increase in the early stages of recovery. Patients with a history of suicide-related events or those exhibiting a significant degree of suicidal thoughts prior to commencement of treatment are known to be at a greater risk of suicidal thoughts or suicidal behaviour, and should receive careful monitoring during treatment. A meta-analysis of placebocontrolled clinical trials of antidepressant medicinal products in psychiatric disorders showed an increased risk of suicidal behaviour with antidepressants compared to placebo in patients less than 25 years old.

Cases of suicidal thoughts and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.8). Physicians should encourage patients to report any distressing thoughts or feelings or depressive symptoms at any time. If while on ARICLAIM therapy, the patient develops agitation or depressive symptoms, specialised medical advice should be sought, as depression is a serious medical condition. If a decision to initiate antidepressant pharmacological therapy is taken, the gradual discontinuation of ARICLAIM is recommended (see Section 4.2).

Use in children and adolescents under 18 years of age

No clinical trials have been conducted with duloxetine in paediatric populations. ARICLAIM should not be used in the treatment of children and adolescents under the age of 18 years. Suicide-related behaviours (suicide attempts and suicidal thoughts), and hostility (predominantly aggression, oppositional behaviour and anger), were more frequently observed in clinical trials among children and adolescents treated with antidepressants compared to those treated with placebo. If, based on clinical need, a decision to treat is nevertheless taken, the patient should be carefully monitored for the appearance of suicidal symptoms. In addition, long-term safety data in children and adolescents concerning growth, maturation and cognitive and behavioural development are lacking.

Haemorrhage

There have been reports of bleeding abnormalities, such as ecchymoses, purpura and gastrointestinal haemorrhage with selective serotonin reuptake inhibitors (SSRIs) and serotonin/noradrenaline reuptake inhibitors (SNRIs). Caution is advised in patients taking anticoagulants and/or medicinal products known to affect platelet function, and in patients with known bleeding tendencies.

Hyponatremia

Hyponatremia has been reported rarely, predominantly in the elderly, when administering ARICLAIM. Caution is required in patients at increased risk for hyponatremia; such as elderly, cirrhotic, or dehydrated patients or patients treated with diuretics. Hyponatremia may be due to a syndrome of inappropriate anti-diuretic hormone secretion (SIADH).

Discontinuation of treatment

Withdrawal symptoms when treatment is discontinued are common, particularly if discontinuation is abrupt (see section 4.8). In clinical trials adverse events seen on abrupt treatment discontinuation occurred in approximately 45% of patients treated with ARICLAIM and 23% of patients taking placebo.

The risk of withdrawal symptoms seen with SSRI's and SNRI's may be dependent on several factors including the duration and dose of therapy and the rate of dose reduction. The most commonly reported reactions are listed in section 4.8. Generally these symptoms are mild to moderate, however, in some patients they may be severe in intensity. They usually occur within the first few days of discontinuing treatment, but there have been very rare reports of such symptoms in patients who have inadvertently missed a dose. Generally these symptoms are self-limiting and usually resolve within 2 weeks, though in some individuals they may be prolonged (2-3 months or more). It is therefore advised that duloxetine should be gradually tapered when discontinuing treatment over a period of no less than 2 weeks, according to the patient's needs (see section 4.2).

Akathisia/psychomotor restlessness

The use of duloxetine has been associated with the development of akathisia, characterised by a subjectively unpleasant or distressing restlessness and need to move often accompanied by an inability to sit or stand still. This is most likely to occur within the first few weeks of treatment. In patients who develop these symptoms, increasing the dose may be detrimental.

Medicinal products containing duloxetine

Duloxetine is used under different trademarks in several indications (treatment of diabetic neuropathic pain, major depressive episodes as well as stress urinary incontinence). The use of more than one of these products concomitantly should be avoided.

Hepatitis/increased liver enzymes

Cases of liver injury, including severe elevations of liver enzymes (>10 times upper limit of normal), hepatitis and jaundice have been reported with duloxetine (see section 4.8). Most of them occurred during the first months of treatment. The pattern of liver damage was predominantly hepatocellular. Duloxetine should be used with caution in patients treated with other medicinal products associated with hepatic injury.

Sucrose

ARICLAIM gastro-resistant capsules contain sucrose. Patients with rare hereditary problems of fructose intolerance, glucose-galactose malabsorption or sucrose-isomaltase insufficiency should not take this medicine.

4.5 Interaction with other medicinal products and other forms of interaction

CNS medicinal products: the risk of using duloxetine in combination with other CNS-active medicinal products has not been systematically evaluated, except in the cases described in this section. Consequently, caution is advised when ARICLAIM is taken in combination with other centrally acting medicinal products and substances including alcohol and sedative medicinal products (e.g. benzodiazepines, morphinomimetics, antipsychotics, phenobarbital, sedative antihistamines).

Monoamine Oxidase Inhibitors (MAOIs): due to the risk of serotonin syndrome, ARICLAIM should not be used in combination with nonselective irreversible monoamine oxidase inhibitors (MAOIs), or within at least 14 days of discontinuing treatment with an MAOI. Based on the half-life of duloxetine, at least 5 days should be allowed after stopping ARICLAIM before starting an MAOI (see section 4.3).

For selective, reversible MAOIs, like moclobemide, the risk of serotonin syndrome is lower. However, the concomitant use of ARICLAIM with selective, reversible MAOIs is not recommended (see section 4.4).

Serotonin syndrome: in rare cases, serotonin syndrome has been reported in patients using SSRIs (e.g.

paroxetine, fluoxetine) concomitantly with serotonergic medicinal products. Caution is advisable if ARICLAIM is used concomitantly with serotonergic antidepressants like SSRIs, tricyclics like clomipramine or amitriptyline, St John's wort (Hypericum perforatum), venlafaxine or triptans, tramadol, pethidine and tryptophan.

Effect of duloxetine on other medicinal products

Medicinal products metabolised by CYP1A2: The pharmacokinetics of theophylline, a CYP1A2 substrate, were not significantly affected by co-administration with duloxetine (60 mg twice daily). Medicinal products metabolised by CYP2D6: Duloxetine is a moderate inhibitor of CYP2D6. When duloxetine was administered at a dose of 60 mg twice daily with a single dose of desipramine, a CYP2D6 substrate, the AUC of desipramine increased 3-fold. The co-administration of duloxetine (40 mg twice daily) increases steady state AUC of tolterodine (2 mg twice daily) by 71 %, but does not affect the pharmacokinetics of its active 5-hydroxyl metabolite and no dosage adjustment is recommended. Caution is advised if ARICLAIM is co-administered with medicinal products that are predominantly metabolised by CYP2D6 (risperidone, tricyclic antidepressants [TCAs] such as nortriptyline, amitriptyline, and imipramine) particularly if they have a narrow therapeutic index (such as flecainide, propafenone and metoprolol).

Oral contraceptives and other steroidal agents: results of *in vitro* studies demonstrate that duloxetine does not induce the catalytic activity of CYP3A. Specific *in vivo* drug interaction studies have not been performed.

Anticoagulants and antiplatelet agents: Caution should be exercised when duloxetine is combined with oral anticoagulants or antiplatelet agents due to a potential increased risk of bleeding. Furthermore, increases in INR values have been reported when duloxetine was co-administered with warfarin.

Effects of other medicinal products on duloxetine

Antacids and H2 antagonists: co-administration of duloxetine with aluminium- and magnesium-containing antacids or duloxetine with famotidine had no significant effect on the rate or extent of duloxetine absorption after administration of a 40 mg oral dose.

Inhibitors of CYP 1A2: because CYP1A2 is involved in duloxetine metabolism, concomitant use of duloxetine with potent inhibitors of CYP1A2 is likely to result in higher concentrations of duloxetine. Fluvoxamine (100 mg once daily), a potent inhibitor of CYP1A2, decreased the apparent plasma clearance of duloxetine by about 77% and increased AUC_{0-t} 6-fold. Therefore ARICLAIM should not be administered in combination with potent inhibitors of CYP1A2 like fluvoxamine (see section 4.3).

Inducers of CYP1A2: Population pharmacokinetic analyses have shown that smokers have almost 50% lower plasma concentrations of duloxetine compared with non-smokers.

4.6 Pregnancy and lactation

Pregnancy

There are no data on the use of duloxetine in pregnant women. Studies in animals have shown reproductive toxicity at systemic exposure levels (AUC) of duloxetine lower than the maximum clinical exposure (see section 5.3).

The potential risk for humans is unknown. As with other serotonergic medicinal products, discontinuation symptoms may occur in the neonate after maternal duloxetine use near term. ARICLAIM should be used in pregnancy only if the potential benefit justifies the potential risk to the foetus. Women should be advised to notify their physician if they become pregnant, or intend to become pregnant, during therapy.

Breast feeding

Duloxetine is very weakly excreted into human milk based on a study of 6 lactating patients, who did not breast feed their children. The estimated daily infant dose on a mg/kg basis is approximately

0.14% of the maternal dose (see section 5.2). As the safety of duloxetine in infants is not known, the use of ARICLAIM while breast-feeding is not recommended.

4.7 Effects on ability to drive and use machines

No studies on the effects on the ability to drive and use machines have been performed. ARICLAIM may be associated with sedation and dizziness. Patients should be instructed that if they experience sedation or dizziness they should avoid potentially hazardous tasks such as driving or operating machinery.

4.8 Undesirable effects

Table 1 gives the adverse reactions observed from spontaneous reporting and in placebo-controlled clinical trials (comprising a total of 6828 patients, 4199 on duloxetine and 2629 on placebo). The most commonly observed adverse reactions in patients with diabetic neuropathic pain treated with ARICLAIM were: nausea; headache, dry mouth and somnolence and dizziness.

Table 1: Adverse reactions

Frequency estimate: Very common ($\geq 1/10$), common ($\geq 1/100$ and <1/10), uncommon ($\geq 1/1,000$ and <1/100), rare ($\geq 1/10,000$ and <1/1,000), very rare (<1/10,000), not known (cannot be estimated from the available data).

Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Investigations					•
	Weight decrease	Weight increase Creatine phosphokinase increased	Blood cholesterol increased		
Cardiac Disorder	S	,			-
	Palpitations	Tachycardia Supra- ventricular arrhythmia, mainly atrial fibrillation			
Nervous System L	Disorders	·			
Headache (14.3%) Somnolence (10.7%) Dizziness (10.2%)	Tremor Paraesthesia	Myoclonus NervousnessDis turbance in attention Lethargy Dysgeusia Dyskinesia Restless legs syndrome Poor quality sleep	Convulsion ¹		Serotonin syndrome Extra-pyramidal symptoms Akathisia Psychomotor restlessness
Eye Disorders			,		
	Blurred vision	Mydriasis Visual disturbance	Glaucoma		

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Ear and Labyrint	h Disorders	I			
	Tinnitus ¹	Vertigo Ear pain			
Respiratory, thore	acic and mediastind				
	Yawning	Throat tightness Epistaxis			
Gastrointestinal I	Disorders				II.
Nausea (24.3%)	Constipation	Gastroenteritis	Stomatitis		Gastrointestina-l
Dry mouth	Diarrhoea	Eructation	Breath odour		haemorrhage
(12.8%)	Vomiting	Gastritis	Haematochezi-a		
	Dyspepsia				
	Flatulence				
Renal and Urinar	y Disorders	1	,		1
		Urinary	Urine odour		
		Retention	abnormal		
		Dysuria			
		Urinary			
		hesitation			
		Nocturia			
		Polyuria Urine flow			
		decreased			
Chin and Subauta	<u> </u> neous Tissue Disor				
Skin ana Subcutai	Sweating	Night sweats			Angio-neurotic
	increased	Urticaria			oedema
	Rash	Dermatitis			Stevens-
	Rasii	contact			Johnson
		Cold sweat			Syndrome
		Photo-			Synarome
		sensitivity			
		reactions			
		Increased			
		tendency to			
		bruise			
Musculoskeletal a	ınd connective tissi	ıe disorders	1		
	Musculo-	Muscle	Trismus		
	skeletal pain	twitching			
	Muscle				
	tightness				
	Muscle spasm				
Endocrine disorde	ers	T			
			Hypo- thyroidism		
Metabolism and N	Nutrition Disorders				
	Decreased	Hyperglycemi-a	Dehydration		SIADH
	Appetite	(reported	Hyponatremia		
		especially in			
		diabetic			
		patients)			
Infections and inf	estations	T	,		
		Laryngitis			

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known
Vascular Disorde	ers		l l		11110 ((11
	Flushing	Blood pressure increase Peripheral coldness Orthostatic hypotension ² Syncope ²			Hypertension Hypertensive crisis
General Disorder	rs and Administratio	on Site Conditions	<u>l</u>		_1
	Fatigue Abdominal pain	Feeling abnormal Feeling cold Thirst Chills Malaise Feeling hot Gait disturbance			Chest pain
Immune system di	isorders				
			Hyper- sensitivity disorder Anaphylactic reaction		
Hepato-biliary di	sorders				
		Elevated liver enzymes (ALT, AST, alkaline phosphatase) Hepatitis ³ Acute liver injury			Jaundice Hepatic failure
Reproductive Syst	l tem and Breast Disc	 orders			
	Erectile dysfunction	Ejaculation disorder Ejaculation delayed Sexual dysfunction Gynaecological haemorrhage	Menopausal symptoms		
Psychiatric Disor					
	Insomnia Agitation Libido decreased Anxiety Orgasm abnormal Abnormal dreams	Sleep disorder Bruxism Disorientation Apathy	Mania Hallucinations Aggression and anger ⁴		Suicidal ideation ⁵ Suicidal ⁵ behaviour

Very common	Common	Uncommon	Rare	Very Rare	Frequency not known

¹ Cases of convulsion and cases of tinnitus have also been reported after treatment discontinuation.

Discontinuation of duloxetine (particularly when abrupt) commonly leads to withdrawal symptoms. Dizziness, sensory disturbances (including paraesthesia), sleep disturbances (including insomnia and intense dreams), agitation or anxiety, nausea and/or vomiting, tremor, headache, irritability, diarrhoea, hyperhydrosis and vertigo are the most commonly reported reactions.

Generally, for SSRIs and SNRIs, these events are mild to moderate and self-limiting, however, in some patients they may be severe and/or prolonged. It is therefore advised that when duloxetine treatment is no longer required, gradual discontinuation by dose tapering should be carried out (see sections 4.2 and 4.4).

In the 12 week acute phase of three clinical trials of duloxetine in patients with diabetic neuropathic pain, small but statistically significant increases in fasting blood glucose were observed in duloxetine-treated patients HbA1c was stable in both duloxetine-treated and placebo-treated patients. In the extension phase of these studies, which lasted up to 52 weeks, there was an increase in HbA1c in both the duloxetine and routine care groups, but the mean increase was 0.3% greater in the duloxetine-treated group. There was also a small increase in fasting blood glucose and in total cholesterol in duloxetine-treated patients while those laboratory tests showed a slight decrease in the routine care group.

Electrocardiograms were obtained from 528 duloxetine-treated and 205 placebo-treated patients with diabetic neuropathic pain in clinical trials lasting up to 13-weeks. The heart rate-corrected QT interval in duloxetine-treated patients did not differ from that seen in placebo-treated patients. No clinically significant differences were observed for QT, PR, QRS, or QTcB measurements between duloxetine-treated and placebo-treated patients.

4.9 Overdose

Cases of overdoses, alone or in combination with other medicinal products, with duloxetine doses of 5400mg were reported. Some fatalities have occurred, primarily with mixed overdoses, but also with duloxetine alone at a dose of approximately 1000 mg. Signs and symptoms of overdose (duloxetine alone or in combination with other medicinal products) included somnolence, coma, serotonin syndrome, seizures, vomiting and tachycardia.

No specific antidote is known for duloxetine but if serotonin syndrome ensues, specific treatment (such as with cyproheptadine and/or temperature control) may be considered. A free airway should be established. Monitoring of cardiac and vital signs is recommended, along with appropriate symptomatic and supportive measures. Gastric lavage may be indicated if performed soon after ingestion or in symptomatic patients. Activated charcoal may be useful in limiting absorption. Duloxetine has a large volume of distribution and forced diuresis, haemoperfusion, and exchange perfusion are unlikely to be beneficial.

²Cases of orthostatic hypotension and syncope have been reported especially at the initiation of treatment.

³See section 4.4

⁴Cases of aggression and anger have been reported particularly early in treatment of after treatment discontinuation.

⁵Cases of suicidal ideation and suicidal behaviours have been reported during duloxetine therapy or early after treatment discontinuation (see section 4.4)

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other antidepressants. ATC code: N06AX21.

Duloxetine is a combined serotonin (5-HT) and noradrenaline (NA) reuptake inhibitor. It weakly inhibits dopamine reuptake with no significant affinity for histaminergic, dopaminergic, cholinergic and adrenergic receptors. Duloxetine dose-dependently increases extracellular levels of serotonin and noradrenaline in various brain areas of animals.

Duloxetine normalised pain thresholds in several preclinical models of neuropathic and inflammatory pain and attenuated pain behaviour in a model of persistent pain. The pain inhibitory action of duloxetine is believed to be a result of potentiation of descending inhibitory pain pathways within the central nervous system.

The efficacy of duloxetine as a treatment for diabetic neuropathic pain was established in 2 randomised, 12-week, double-blind, placebo-controlled, fixed dose studies in adults (22 to 88 years) having diabetic neuropathic pain for at least 6 months. Patients meeting diagnostic criteria for major depressive disorder were excluded from these trials. The primary outcome measure was the weekly mean of 24-hour average pain, which was collected in a daily diary by patients on an 11-point Likert scale.

In both studies, duloxetine 60 mg once daily and 60 mg twice daily significantly reduced pain compared with placebo. The effect in some patients was apparent in the first week of treatment. The difference in mean improvement between the two active treatment arms was not significant. At least 30% reported pain reduction was recorded in approximately 65% of duloxetine treated patients versus 40% for placebo. The corresponding figures for at least 50% pain reduction were 50% and 26% respectively. Clinical response rates (50% or greater improvement in pain) were analysed according to whether or not the patient experienced somnolence during treatment. For patients not experiencing somnolence, clinical response was observed in 47% of patients receiving duloxetine and 27% patients on placebo. Clinical response rates in patients experiencing somnolence were 60% on duloxetine and 30% on placebo. Patients not demonstrating a pain reduction of 30% within 60 days of treatment were unlikely to reach this level during further treatment

In an open label long-term uncontrolled study, the pain reduction in patients responding to 8-weeks of acute treatment of ARICLAIM 60 mg once daily was maintained for a further 6-months as measured by change on the Brief Pain Inventory(BPI) 24-hour average pain item.

5.2 Pharmacokinetic properties

Duloxetine is administered as a single enantiomer. Duloxetine is extensively metabolised by oxidative enzymes (CYP1A2 and the polymorphic CYP2D6), followed by conjugation. The pharmacokinetics of duloxetine demonstrate large intersubject variability (generally 50-60%), partly due to gender, age, smoking status and CYP2D6 metaboliser status.

Duloxetine is well absorbed after oral administration with a Cmax occurring 6 hours post dose. The absolute oral bioavailability of duloxetine ranged from 32% to 80% (mean of 50%). Food delays the time to reach the peak concentration from 6 to 10 hours and it marginally decreases the extent of absorption (approximately 11 %). These changes do not have any clinical significance. Duloxetine is approximately 96% bound to human plasma proteins. Duloxetine binds to both albumin and alphalacid glycoprotein. Protein binding is not affected by renal or hepatic impairment.

Duloxetine is extensively metabolised and the metabolites are excreted principally in urine. Both cytochromes P450-2D6 and 1A2 catalyse the formation of the two major metabolites glucuronide conjugate of 4-hydroxy duloxetine and sulphate conjugate of 5hydroxy,6-methoxy duloxetine. Based upon *in vitro* studies, the circulating metabolites of duloxetine are considered pharmacologically inactive. The pharmacokinetics of duloxetine in patients who are poor metabolisers with respect to

CYP2D6 has not been specifically investigated. Limited data suggest that the plasma levels of duloxetine are higher in these patients.

The elimination half-life of duloxetine ranges from 8 to 17 hours (mean of 12 hours). After an intravenous dose the plasma clearance of duloxetine ranges from 22 l/hr to 46 l/hr (mean of 36 l/hr). After an oral dose the apparent plasma clearance of duloxetine ranges from 33 to 261 l/hr (mean 101 l/hr).

Special populations:

Gender: pharmacokinetic differences have been identified between males and females (apparent plasma clearance is approximatively 50% lower in females). Based upon the overlap in the range of clearance, gender-based pharmacokinetic differences do not justify the recommendation for using a lower dose for female patients.

Age: pharmacokinetic differences have been identified between younger and elderly females (\geq 65 years) (AUC increases by about 25% and half-life is about 25% longer in the elderly), although the magnitude of these changes is not sufficient to justify adjustments to the dose. As a general recommendation, caution should be exercised when treating the elderly (see sections 4.2 and 4.4).

Renal impairment: end stage renal disease (ESRD) patients receiving dialysis had 2-fold higher duloxetine Cmax and AUC values compared with healthy subjects. Pharmacokinetic data on duloxetine is limited in patients with mild or moderate renal impairment.

Hepatic impairment: moderate liver disease (Child Pugh Class B) affected the pharmacokinetics of duloxetine. Compared with healthy subjects, the apparent plasma clearance of duloxetine was 79% lower, the apparent terminal half-life was 2.3 times longer, and the AUC was 3.7 times higher in patients with moderate liver disease. The pharmacokinetics of duloxetine and its metabolites have not been studied in patients with mild or severe hepatic insufficiency.

Nursing mothers: The disposition of duloxetine was studied in 6 lactating women who were at least 12-weeks postpartum. Duloxetine is detected in breast milk, and steady-state concentrations in breast milk are about one-fourth those in plasma. The amount of duloxetine in breast milk is approximately $7~\mu g/day$ while on 40 mg twice daily dosing. Lactation did not influence duloxetine pharmacokinetics

5.3 Preclinical safety data

Duloxetine was not genotoxic in a standard battery of tests and was not carcinogenic in rats. Multinucleated cells were seen in the liver in the absence of other histopathological changes in the rat carcinogenicity study. The underlying mechanism and the clinical relevance are unknown. Female mice receiving duloxetine for 2 years had an increased incidence of hepatocellular adenomas and carcinomas at the high dose only (144 mg/kg/day), but these were considered to be secondary to hepatic microsomal enzyme induction. The relevance of this mouse data to humans is unknown. Female rats receiving duloxetine (45 mg/kg/day) before and during mating and early pregnancy had a decrease in maternal food consumption and body weight, oestrous cycle disruption, decreased live birth indices and progeny survival, and progeny growth retardation at systemic exposure levels estimated to be at the most at maximum clinical exposure (AUC). In an embryotoxicity study in the rabbit, a higher incidence of cardiovascular and skeletal malformations was observed at systemic exposure levels below the maximum clinical exposure (AUC). No malformations were observed in another study testing a higher dose of a different salt of duloxetine. In prenatal/postnatal toxicity studies in the rat, duloxetine induced adverse behavioural effects in the offspring at exposures below maximum clinical exposure (AUC).

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Capsule content:

Hypromellose.

Hypromellose acetate succinate

Sucrose

Sugar spheres

Talc

Titanium dioxide (E171)

Triethyl citrate.

Capsule shell:

60 mg:

Gelatin

Sodium Lauryl Sulfate

Titanium Dioxide (E171)

Indigo Carmine (E132)

Yellow Iron Oxide (E172)

Edible White Ink

Edible White Ink contains:

Titanium Dioxide (E171)

Propylene glycol

Shellac

Povidone.

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years.

6.4 Special precautions for storage

Store in the original package in order to protect from moisture. Do not store above 30 C.

6.5 Nature and contents of container

Polyvinylchloride (PVC), Polyethylene (PE), and Polychlorotrifluoroethylene (PCTFE) blister sealed with an aluminum foil.

ARICLAIM 60 mg is available in packs of 28 and 98 capsules.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

7. MARKETING AUTHORISATION HOLDER

Eli Lilly Nederland BV, Grootslag 1-5, NL-3991 RA Houten, The Netherlands.

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/04/283/011 EU/1/04/283/012

- 9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION
- 10. DATE OF REVISION OF THE TEXT

ANNEX II

- A. MANUFACTURING AUTHORISATION HOLDER(S) RESPONSIBLE FOR BATCH RELEASE
- B. CONDITIONS OF THE MARKETING AUTHORISATION

A MANUFACTURING AUTHORISATION HOLDER(S) RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer(s) responsible for batch release

Lilly S.A. Avda. de la Industria Nº 30, 28108 Alcobendas Madrid Spain

- **B** CONDITIONS OF THE MARKETING AUTHORISATION
- CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE IMPOSED ON THE MARKETING AUTHORISATION HOLDER

Medicinal product subject to medical prescription

• CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

Not applicable.

• OTHER CONDITIONS

ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
CARTONS FOR 40 MG HARD GASTRO-RESISTANT CAPSULES
1. NAME OF THE MEDICINAL PRODUCT
ARICLAIM 40 mg, hard gastro-resistant capsules. Duloxetine
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each capsule contains 40 mg duloxetine (as hydrochloride)
3. LIST OF EXCIPIENTS
Contains sucrose See leaflet for further information
4. PHARMACEUTICAL FORM AND CONTENTS
28 hard gastro-resistant capsules 56 hard gastro-resistant capsules 98 hard gastro-resistant capsules 140 hard gastro-resistant capsules
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Oral use. Read the package leaflet before use.
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE REACH AND SIGHT OF CHILDREN
Keep out of the reach and sight of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
EXP {MM/YYYY}.

Store in the original package. Do not store above 30 °C.
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Eli Lilly Nederland BV, Grootslag 1-5, NL-3991 RA, Houten, The Netherlands.
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/04/283/002-005
13. BATCH NUMBER
Lot:
14. GENERAL CLASSIFICATION FOR SUPPLY
Medicinal product subject to medical prescription.
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
ARICLAIM 40 mg

9.

SPECIAL STORAGE CONDITIONS

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

CARTON FOR 98 CAPSULES (40 MG) AS INTERMEDIATE PACK / COMPONENT OF A MULTIPACK (WITHOUT BLUE BOX)

1. NAME OF THE MEDICINAL PRODUCT

ARICLAIM 40 mg hard gastro-resistant capsules Duloxetine

2. STATEMENT OF ACTIVE SUBSTANCE(S)

Each capsule contains 40 mg duloxetine (as hydrochloride).

3. LIST OF EXCIPIENTS

Contains sucrose.

See leaflet for further information

4. PHARMACEUTICAL FORM AND CONTENTS

98 capsules

Component of a multipack comprising 2 packs, each containing 98 capsules.

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Oral use.

Read the package leaflet before use.

6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE REACH AND SIGHT OF CHILDREN

Keep out of the reach and sight of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8. EXPIRY DATE

EXP {MM/YYYY}

9. SPECIAL STORAGE CONDITIONS

Store in the original package. Do not store above 30 °C.

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF
APPROPRIATE
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Eli Lilly Nederland BV, Grootslag 1-5, NL-3991 RA, Houten, The Netherlands.
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/04/283/004
13. BATCH NUMBER
Lot:
14. GENERAL CLASSIFICATION FOR SUPPLY
Medicinal product subject to medical prescription.
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
ARICLAIM 40 mg

PARTICULARS TO APPEAR ON THE OUTER PACKAGING OUTER WRAPPER LABEL ON MULTIPACKS (2X98 CAPSULES, 40 MG) WRAPPED IN FOIL (INCLUDING BLUE BOX)

1. NAME OF THE MEDICINAL PRODUCT

ARICLAIM 40 mg hard gastro-resistant capsules Duloxetine

2. STATEMENT OF ACTIVE SUBSTANCE(S)

Each capsule contains 40 mg duloxetine (as hydrochloride).

3. LIST OF EXCIPIENTS

Contains sucrose

See leaflet for further information

4. PHARMACEUTICAL FORM AND CONTENTS

Multipack comprising 2 packs, each containing 98 capsules.

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Oral use

Read the package leaflet before use.

6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE REACH AND SIGHT OF CHILDREN

Keep out of the reach and sight of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8. EXPIRY DATE

EXP {MM/YYYY}

9. SPECIAL STORAGE CONDITIONS

Store in the original package. Do not store above 30 °C.

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF

	APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Eli L	illy Nederland BV, Grootslag 1-5, NL-3991 RA, Houten, The Netherlands.
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1	/04/283/006
13.	BATCH NUMBER
Lot:	
14.	GENERAL CLASSIFICATION FOR SUPPLY
Medi	cinal product subject to medical prescription.
15.	INSTRUCTIONS ON USE
16	DECORMATION BY DRAW I.E.
16.	INFORMATION IN BRAILLE

ARICLAIM 40 mg

resist	ant capsules)
1.	NAME OF THE MEDICINAL PRODUCT
ARIC	LAIM 40 mg hard gastro-resistant capsules
Dulox	
_ ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
2.	NAME OF THE MARKETING AUTHORISATION HOLDER
Lilly	
3.	EXPIRY DATE
<exf< td=""><td>P {MM/YYYY}.</td></exf<>	P {MM/YYYY}.
4.	BATCH NUMBER
Lot:	
5.	OTHER

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS (40 mg hard gastro-

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
CARTONS FOR 20 MG HARD GASTRO-RESISTANT CAPSULES
1. NAME OF THE MEDICINAL PRODUCT
ARICLAIM 20 mg, hard gastro-resistant capsules. Duloxetine
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each capsule contains 20 mg duloxetine (as hydrochloride).
3. LIST OF EXCIPIENTS
Contains sucrose. See leaflet for further information
4. PHARMACEUTICAL FORM AND CONTENTS
28 hard gastro-resistant capsules 56 hard gastro-resistant capsules.
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Oral use. Read the leaflet before use
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE REACH AND SIGHT OF CHILDREN
Keep out of the reach and sight of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
EXP {MM/YYYY}.
9. SPECIAL STORAGE CONDITIONS
Store in the original package. Do not store above 30 °C. 10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF

APPROPRIATE

12.	MARKETING AUTHORISATION NUMBER(S)
EU/	1/04/283/001
	1/04/283/007
13.	BATCH NUMBER
Lot:	
14.	GENERAL CLASSIFICATION FOR SUPPLY
	GENERAL CLASSIFICATION FOR SUPPLY dicinal product subject to medical prescription.
Med	
14. Med	licinal product subject to medical prescription.

NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

11.

resistant capsules)	
1.	NAME OF THE MEDICINAL PRODUCT
ARIC	LAIM 20 mg hard gastro-resistant capsules
Dulox	retine
2.	NAME OF THE MARKETING AUTHORISATION HOLDER
Lilly	
	EXPLOYED A TELE
3.	EXPIRY DATE
∠E3/D	
<exp< td=""><td>{MM/YYYY}.</td></exp<>	{MM/YYYY}.
4	RATCH NUMRER
т.	DATCH NUMBER
Lot:	
5.	OTHER
4. Lot:	OTHER

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS (20 mg hard gastro-

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
CARTONS FOR 30 MG HARD GASTRO-RESISTANT CAPSULES
1. NAME OF THE MEDICINAL PRODUCT
ARICLAIM 30 mg, hard gastro-resistant capsules. Duloxetine
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each capsule contains 30 mg of duloxetine (as hydrochloride)
3. LIST OF EXCIPIENTS
Contains sucrose See leaflet for further information
4. PHARMACEUTICAL FORM AND CONTENTS
7 hard gastro-resistant capsules, 28 hard gastro-resistant capsules 98 hard gastro-resistant capsules
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Oral use. Read the package leaflet before use.
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE REACH AND SIGHT OF CHILDREN
Keep out of the reach and sight of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
EXP
9. SPECIAL STORAGE CONDITIONS

Store in the original package. Do not store above 30°C

10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Eli Li	lly Nederland BV, Grootslag 1-5, NL-3991 RA, Houten, The Netherlands.
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1/	/04/283/008 /04/283/009 /04/283/010
13.	BATCH NUMBER
Lot:	
14.	GENERAL CLASSIFICATION FOR SUPPLY
	cinal product subject to medical prescription.
15.	INSTRUCTIONS ON USE
-	
16.	INFORMATION IN BRAILLE
ARIC	CLAIM 30 mg

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS 30 mg hard gastro-resistant capsules		
1. NAME OF THE MEDICINAL PRODUCT		
ARICLAIM 30 mg hard gastro-resistant capsules Duloxetine		
2. NAME OF THE MARKETING AUTHORISATION HOLDER		
Lilly		
3. EXPIRY DATE		
EXP		
4. BATCH NUMBER		
Lot:		

5.

OTHER

PARTICULARS TO APPEAR ON THE OUTER PACKAGING	
CARTONS FOR 60 MG HARD GASTRO-RESISTANT CAPSULES	
1. NAME OF THE MEDICINAL PRODUCT	
ARICLAIM 60 mg, hard gastro-resistant capsules. Duloxetine	
2. STATEMENT OF ACTIVE SUBSTANCE(S)	
Each capsule contains 60 mg of duloxetine (as hydrochloride)	
3. LIST OF EXCIPIENTS	
Contains sucrose See leaflet for further information	
4. PHARMACEUTICAL FORM AND CONTENTS	
28, hard gastro-resistant capsules 98, hard gastro-resistant capsules.	
5. METHOD AND ROUTE(S) OF ADMINISTRATION	
Oral use. Read the leaflet before use	
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE REACH AND SIGHT OF CHILDREN	
Keep out of the reach and sight of children.	
7. OTHER SPECIAL WARNING(S), IF NECESSARY	
0 EVDIDV DATE	
8. EXPIRY DATE	
EXP	
9. SPECIAL STORAGE CONDITIONS	
Store in the original package. Do not store above 30°C	

10.

SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS

APPROPRIATE
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Eli Lilly Nederland BV, Grootslag 1-5, NL-3991 RA, Houten, The Netherlands.
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/04/283/011
EU/1/04/283/012
13. BATCH NUMBER
Lot:
14. GENERAL CLASSIFICATION FOR SUPPLY
Medicinal product subject to medical prescription.
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
ARICLAIM 60 mg

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS 60 mg hard gastro-resistant capsules		
1. NAME OF THE MEDICINAL PRODUCT		
ARICLAIM 60 mg hard gastro-resistant capsules Duloxetine		
2. NAME OF THE MARKETING AUTHORISATION HOLDER		
Lilly		
3. EXPIRY DATE		
EXP		
4. BATCH NUMBER		
Lot:		

5.

OTHER

B. PACKAGE LEAFLET

PACKAGE LEAFLET: INFORMATION FOR THE USER

ARICLAIM 40 mg hard gastro-resistant capsules ARICLAIM 20 mg hard gastro-resistant capsules

Duloxetine (as hydrochloride)

Read all of this leaflet carefully before you start taking this medicine.

- Keep this leaflet. You may need to read it again
- If you have any further questions, ask your doctor or pharmacist
- This medicine has been prescribed for you. Do not pass it on toothers. It may harm them, even if their symptoms are the same as yours
- If any of the side effects gets serious, or if you notice any side effects not listed in this leaflet, please tell your doctor or pharmacist

In this leaflet:

- 1. What ARICLAIM is and what it is used for
- 2. Before you take ARICLAIM
- 3. How to take ARICLAIM
- 4. Possible side effects
- 5. How to store ARICLAIM
- 6. Further information

1. WHAT ARICLAIM IS AND WHAT IT IS USED FOR

ARICLAIM increases the levels of serotonin and noradrenaline in the nervous system. ARICLAIM is a medicine to be taken by mouth to treat Stress Urinary Incontinence (SUI) in women or to treat a condition called diabetic neuropathic pain in adults.

Stress urinary incontinence is a medical condition in which patients have accidental loss or leakage of urine during physical exertion or activities such as laughing, coughing, sneezing, lifting, or exercise.

ARICLAIM is believed to work by increasing the strength of the muscle that holds back urine when you laugh, sneeze, or perform physical activities.

The efficacy of ARICLAIM is reinforced when combined with a training program called Pelvic Floor Muscle Training (PFMT).

Neuropathic pain is a medical condition in which the pain is commonly described as burning, stabbing, stinging, shooting or aching or like an electric shock. There may be loss of feeling in the affected area, or sensations such as touch, heat, cold or pressure may cause pain.

The effect of ARICLAIM may be noticeable in many patients with diabetic neuropathic pain within 1 week of treatment.

2. BEFORE YOU TAKE ARICLAIM

DO NOT take ARICLAIM if you:

- are allergic (hypersensitive) to duloxetine or any of the inactive ingredients of ARICLAIM
- have liver disease
- have severe kidney disease
- are taking or have taken within the last 14 days, another medicine known as a monoamine oxidase inhibitor MAOI (see also below in section 'Taking other medicines')
- If you are taking fluvoxamine which is usually used to treat depression, ciprofloxacin or enoxacine which are used to treat some infections

Talk to your doctor if you have high blood pressure. Your doctor will tell you if you should be taking ARICLAIM.

Take special care with ARICLAIM

The following are reasons why ARICLAIM may not be suitable for you. Talk to your doctor before you take the medicine if you:

- are taking other medicines to treat depression (see 'Taking other medicines')
- are taking St. John's Wort, a herbal treatment (Hypericum perforatum)
- have kidney disease
- have had seizures (fits)
- have had mania
- suffer from bipolar disorder
- have eye problems, such as certain kinds of glaucoma (increased pressure in the eye)
- have a history of bleeding disorders (tendency to develop bruises)
- are at risk of low sodium levels
- are currently being treated with another medicine which may cause liver damage
- are taking other medicines containing duloxetine
- have intolerance to some sugars (see end of Section 2)
- are considering stopping ARICLAIM (see section 3)

ARICLAIM may cause a sensation of restlessness or an inability to sit or stand still. You should tell your doctor if this happens to you.

Thoughts of suicide and worsening of your depression or anxiety disorder

If you are depressed and/or have anxiety disorders you can sometimes have thoughts of harming or killing yourself. These may be increased when first starting antidepressants, since these medicines all take time to work, usually about two weeks but sometimes longer.

You may be more likely to think like this if you:

- have previously had thoughts about killing or harming yourself.
- are a young adult. Information from clinical trials has shown an increased risk of suicidal behaviour in adults aged less than 25 years with psychiatric conditions who were treated with an antidepressant.

If you_have thoughts of harming or killing yourself at any time, contact your doctor or go to a hospital straight away.

You may find it helpful to tell a relative or close friend that you are depressed or have_an anxiety disorder, and ask them to read this leaflet. You might ask them to tell you if they think your depression or anxiety is getting worse, or if they are worried about changes in your behaviour.

Use in children and adolescents under 18 years of age

ARICLAIM should not be used for children and adolescents under the age of 18 years. Also, you should know that patients under 18 have an increased risk of side-effects such as suicide attempt, suicidal thoughts and hostility (predominantly aggression, oppositional behaviour and anger) when they take this class of medicines. Also, the long-term safety effects concerning growth, maturation, and cognitive and behavioural development of ARICLAIM in this age group have not yet been demonstrated.

Taking other medicines

Please tell your doctor or pharmacist if you are taking or have recently taken any other medicines, including medicines obtained without a prescription. The main ingredient of ARICLAIM, duloxetine, is used in other medicines for other conditions:

• depression, anxiety and urinary incontinence

Using more than one of these medicines at the same time should be avoided. Check with your doctor if you are already taking other medicines containing duloxetine.

Your doctor should decide whether you can take ARICLAIM with other medicines. Do not start or stop taking any medicines, including those bought without a prescription and herbal remedies, before checking with your doctor.

You should also tell your doctor if you are taking any of the following:

Monoamine Oxidase Inhibitors (MAOI): you should not take ARICLAIM with an MAOI or within 14 days of stopping an MAOI. Taking an MAOI together with many prescription medicines, including ARICLAIM, can cause serious or even life-threatening side effects. You must wait at least 14 days after you have stopped taking an MAOI before you can take ARICLAIM. Also, you need to wait at least 5 days after you stop taking ARICLAIM before you take an MAOI.

Medicines that cause sleepiness: These include medicines prescribed by your doctor including alcohol and sedative medicines (benzodiazepines, strong painkillers, antipsychotics, phenobarbital, sedative antihistamines).

Medicines that increase the level of serotonin: triptans, tramadol, tryptophan, SSRIs (such as paroxetine and fluoxetine), tricyclics (such as clomipramine, amitriptyline) and venlafaxine. These medicines increase the risk of side effects; if you get any unusual symptom taking any of these medicines together with ARICLAIM, you should see your doctor.

Oral-anticoagulants: medicines which thin the blood. These medicines might increase the risk of bleeding.

Taking ARICLAIM with food and drink

ARICLAIM may be taken with or without food. You should take extra care if you drink alcohol while taking ARICLAIM.

Pregnancy and breast-feeding

Ask your doctor or pharmacist for advice before taking any medicine.

Tell your doctor if you:

become pregnant, or you are trying to become pregnant, while you are taking ARICLAIM. You should use ARICLAIM only after discussing the potential benefits and any potential risks to your unborn child with your doctor.

• breast-feeding. The use of ARICLAIM while breastfeeding is not recommended. You should ask your doctor or pharmacist for advice.

Driving and using machines

ARICLAIM may make you feel sleepy of dizzy. Do not drive or use any tools or machines until you know how ARICLAIM affects you.

Important information about some of the ingredients of ARICLAIM

ARICLAIM contains **sucrose.** If you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this medicinal product.

3. HOW TO TAKE ARICLAIM

Always take ARICLAIM exactly as your doctor has told you. You should check with your doctor or pharmacist if you are not sure.

The recommended dose of ARICLAIM for the treatment of Stress Urinary Incontinence is 40 mg twice a day (in the morning and late afternoon/evening). Your doctor may decide to start your treatment with one capsule of 20 mg twice a day for two weeks before increasing the dose to 40 mg twice a day.

The usual dose of ARICLAIM for the treatment Neuropathic pain is 60 mg once a day, but your doctor will prescribe the dose that is right for you.

ARICLAIM is for oral use. You should swallow your capsule whole with a drink of water.

To help you remember to take ARICLAIM, you may find it easier to take it at the same times every day.

Do not stop taking ARICLAIM without talking to your doctor.

If you take more ARICLAIM than you should

Call your doctor or pharmacist immediately if you take more than the amount of ARICLAIM prescribed by your doctor.

If you forget to take ARICLAIM

If you miss a dose, take it as soon as you remember. However, if it is time for your next dose, skip the missed dose and take only a single dose as usual. Do not take a double dose to make up for forgotten doses. Do not take more than the daily amount of ARICLAIM that has been prescribed for you in one day.

If you stop taking ARICLAIM

DO NOT stop taking your capsules without the advice of your doctor even if you feel better. If your doctor thinks that you no longer need ARICLAIM he or she will ask you to reduce your dose over 2 weeks.

Some patients, who suddenly stop taking ARICLAIM after more than 1 week of therapy, have had symptoms such as:

• dizziness, tingling feelings like pins and needles, sleep disturbances (vivid dreams, nightmares, inability to sleep), feeling restless or agitated, feeling anxious, feeling sick (nausea) or being sick (vomiting), tremor (shakiness), headaches, feeling irritable, diarrhoea, excessive sweating or vertigo. These symptoms are usually not serious and disappear within a few days, but if you have symptoms that are troublesome you should ask your doctor for advice.

If you have any further questions on the use of this product, ask your doctor or pharmacist.

4. POSSIBLE SIDE EFFECTS

Like all medicines, ARICLAIM can cause side effects, although not everybody gets them. These effects are normally mild to moderate and often disappear after a short time.

Very common side effects (these can affect more than 1 in 10 patients treated) with ARICLAIM when taken to treat stress urinary incontinence

• feeling sick (nausea), dry mouth, constipation and tiredness

Very common side effects (these can affect more than 1 in 10 patients treated) with ARICLAIM when taken to treat diabetic neuropathic pain

• feeling sick (nausea), somnolence, headache, dizziness and dry mouth

Common side effects (these can affect from 1 to 10 users in 100 patients treated)

- anxiety, feeling agitated, less sex drive, having a poor quality of sleep
- headache, dizziness, feeling sleepy, tremor or numbness, including numbness or tingling of the skin
- flushing
- stomach pain, shivering, weakness
- being sick (vomiting), diarrhoea, heartburn
- increased sweating
- lack of appetite
- blurred eyesight
- vertigo

Uncommon side effects (these can affect from 1 to 10 users in 1,000 patients treated)

- throat inflammation
- feeling disorientated, having abnormal dreams, lack of motivation, , grinding of teeth, changes in orgasm
- increased yawning
- tasting things differently than usual, nervousness, disturbance in attention
- muscle pain, muscle tightness or muscle spasm, contraction of the jaw muscle
- weight loss or weight gain, increased level of cholesterol in the blood
- burping, breaking wind, bad breath, indigestion or gastroenteritis
- inflammation of the liver that may cause abdominal pain, tiredness or yellow coloration of the skin
- tinnitus (perception of sound in the ear when there is no external sound), ear pain
- larger pupils (the dark centre of the eye) or visual disturbance
- feeling the heart pumping in the chest, fast or irregular heart beat
- fainting, increase in blood pressure
- increased tendency to bruise, night sweats, cold sweats
- menopausal symptoms, abnormal periods, including heavy or prolonged periods
- allergic reactions, blisters
- decreased thyroid gland activity
- need to pass urine during the night, difficulty or inability to pass urine, pain on passing urine or abnormal urine odour
- dehydration,
- feeling hot/cold, thirst
- (itchy) rash

Rare side effects (these can affect from 1 to 10 users in 10,000 patients treated)

- increased level of sugar in the blood, low levels of sodium in the blood (the symptoms are feeling sick and unwell with weak muscles or confused)
- involuntary movements of the muscles, restless legs syndrome
- muscle twitching
- throat tightness or nose bleeds
- increased pressure in the eye (glaucoma)
- sensitivity to sunlight
- feeling dizzy (particularly when standing up too quickly), feeling cold in your fingers and/or toes
- abnormal manner of walking
- need to pass more urine then usual or urine flow decreased
- serious allergic reaction which causes difficulty in breathing or dizziness or hives
- experiencing hallucinations

Other possible side effects

- mania (over activity, racing thoughts and decrease need for sleep), experiencing aggression and anger, suicidal thoughts or behavioura sensation of restlessness or an inability to sit or stand still, "Serotonin syndrome" (a rare reaction which may cause feelings of great happiness, drowsiness, clumsiness, restlessness, feeling of being drunk, fever, sweating or rigid muscles), fits or stiffness
- passing bright red blood in your stools vomiting blood, or black tarry stools (faeces)
- syndrome of inadequate secretion of anti-diuretic hormone (SIADH)
- chest pain
- yellow colouration of the skin (jaundice), hepatic failure, Stevens-Johnson syndrome, sudden swelling of skin of mucosa (angioedema)

If any of the side effects get serious, or if you notice any side effects not listed in this leaflet, please tell your doctor.

HOW TO STORE ARICLAIM 5.

Keep out of the reach and sight of children

Do not use ARICLAIM after the expiry date which is stated on the carton.

Store in the original pack. Do not store above 30 °C.

6. **FURTHER INFORMATION**

What ARICLAIM contains

The **active** substance is duloxetine.

Each capsule contains 20 or 40 mg of duloxetine (as hydrochloride).

The **other** ingredients are:

Capsule content: hypromellose, hypromellose acetate succinate, sucrose, sugar spheres, talc, titanium dioxide (E171), triethyl citrate.

(See end of section 2 for further information on sucrose)

Capsule shell: gelatin, sodium lauryl sulphate, titanium dioxide (E171), indigo carmine (E132), iron oxide red and iron oxide yellow, edible black ink.

Edible Ink: Black Iron Oxide-Synthetic (E172), Propylene glycol, Shellac.

What ARICLAIM looks like and contents of the pack

ARICLAIM is a hard gastro-resistant capsule.

Each capsule of ARICLAIM contains pellets of the active substance with a covering to protect them from stomach acid.

ARICLAIM is available in 2 strengths: 20 and 40 mg.

The 40 mg capsule has an opaque orange body imprinted with '40 mg' and an opaque blue cap, imprinted with '9545'.

The 20 mg capsule has an opaque blue body imprinted with '20 mg' and an opaque blue cap, imprinted with '9544'.

ARICLAIM 40 mg is available in blister packs of 28, 56, 98, 140 and 196 (2 x 98) capsules. ARICLAIM 20 mg is available in blister packs of 28 and 56 capsules.

Not all pack sizes may be marketed.

Marketing Authorisation Holder and Manufacturer

Marketing Authorisation Holder: Marketing Authorisation Holder: Eli Lilly Nederland BV, Grootslag 1-5, NL-3991 RA Houten, The Netherlands.

Manufacturer: Lilly S.A., Avda. De la Industria, 30, 28108 Alcobendas, Madrid, Spain.

For any information about this medicine product, please contact the local representative of the Marketing Authorisation Holder.

België/Belgique/Belgien

S.C.S. Boehringer Ingelheim Comm.V.

Tél/Tel: +32 27 73 33 11

България

ТП "Ели Лили Недерланд" Б.В. - България тел. + 359 2 491 41 40

Česká republika

Luxembourg/Luxemburg

S.C.S. Boehringer Ingelheim Comm.V.

Tél/Tel: +32 2 773 33 11

Magyarország

Boehringer Ingelheim Pharma

Tel.: +36 1 224 7120

Malta

Boehringer Ingelheim spol. s r.o.

Tel.: + 42 02 34 65 51 11

Danmark

Boehringer Ingelheim Danmark A/S

Tlf: +45 39 15 88 88

Deutschland

Boehringer Ingelheim Pharma GmbH & Co. KG

Tel: +49 (0) 69 50 50 83 09

Eesti

Boehringer Ingelheim Pharma GmbH

Tel: + 37 2 60 80 940

Ελλάδα

ΦΑΡΜΑΣΕΡΒ-ΛΙΛΛΥ Α.Ε.Β.Ε.

Τηλ: +30 210 629 4600

EspañaDista S.A..

Tel: + 34 91 623 17 32

France

Boehringer Ingelheim France S.A.S.

Tél: +33 3 26 50 45 33

Ireland

Boehringer Ingelheim Ireland Ltd.

Tel: +353-(0) 1 661 4377

Ísland

Eli Lilly Danmark A/S, Útibú á Íslandi

Tel: + 354 520 34 00

Italia

Eli Lilly Italia S.p.A.

Tel: +39-055 42571

Κύπρος

Boehringer Ingelheim Ellas A.E.

Τηλ: +30 2 10 89 06 300

Latvija

Boehringer Ingelheim Pharma GmbH

Tel: +37 167 24 00 68

Lietuva

Boehringer Ingelheim Pharma Ges mbH

Tel.: +370 37 47 39 22

Boehringer Ingelheim Ltd. Tel: +356 25600 500

Nederland

Boehringer Ingelheim b.v.

Tel: +31 30 6 02 59 14

Norge

Boehringer Ingelheim Norway KS

Tlf: +47 66 76 13 00

Österreich

Boehringer Ingelheim Austria GmbH

Tel: +43 1 710 3739

Polska

Boehringer Ingelheim Sp.z o.o.

Tel.: +48 22 699 0 699

Portugal

Boehringer Ingelheim, Lda

Tel: +351 21 313 53 00

România

Eli Lilly România S.R.L.

Tel: +40 21 4023000

Slovenija

Boehringer Ingelheim Pharma

Tel.: +386 1 586 40 00

Slovenská republika

Boehringer Ingelheim Pharma

Tel.: +421 2 5341 8445

Suomi/Finland

Oy Eli Lilly Finland Ab

Puh/Tel: +358 9 8545 250

Sverige

Boehringer Ingelheim AB

Tel: +46 8 721 21 00

United Kingdom

Boehringer Ingelheim Ltd.

Tel: +44 (0) 1256 315999

This leaflet was last approved in

Detailed information on this medicine is available on the European Medicines Agency (EMEA) web site: http://www.emea.europa.eu.

PACKAGE LEAFLET: INFORMATION FOR THE USER

ARICLAIM 30 mg hard gastro-resistant capsules ARICLAIM 60 mg hard gastro-resistant capsules

Duloxetine (as hydrochloride)

Read all of this leaflet carefully before you start taking this medicine.

- Keep this leaflet. You may need to read it again
- If you have any further questions, ask your doctor or pharmacist
- This medicine has been prescribed for you. Do not pass it on to others. It may harm them, even if their symptoms are the same as yours
- If any of the side effects gets serious, or if you notice any side effects not listed in this leaflet, please tell your doctor or pharmacist

In this leaflet:

- 1. What ARICLAIM is and what it is used for
- 2. Before you take ARICLAIM
- 3. How to take ARICLAIM
- 4. Possible side effects
- 5 How to store ARICLAIM
- 6. Further information

1. WHAT ARICLAIM IS AND WHAT IT IS USED FOR

ARICLAIM increases the levels of serotonin and noradrenaline in the nervous system.

ARICLAIM is used in adults to treat a condition called diabetic neuropathic pain (often described as burning, stabbing, stinging, shooting or aching or like an electric shock. There may be loss of feeling in the affected area, or sensations such as touch, heat, cold or pressure may cause pain).

The effect of ARICLAIM may be noticeable in many patients with diabetic neuropathic pain within 1 week of treatment.

2. BEFORE YOU TAKE ARICLAIM

DO NOT take ARICLAIM if you:

- are allergic (hypersensitive) to duloxetine or any of the other ingredients of ARICLAIM
- are taking or have recently taken within the last 14 days, another antidepressant medicine called a monoamine oxidase inhibitor (MAOI) (see also below in section: 'Taking other medicines')
- have liver disease
- have severe kidney disease
- are taking fluvoxamine which is usually used to treat depression, ciprofloxacin or enoxacine which are used to treat some infections

Talk to your doctor if you have high blood pressure. Your doctor will tell you if you should be taking ARICLAIM.

Take special care with ARICLAIM

The following are reasons why ARICLAIM may not be suitable for you. Talk to your doctor before you take the medicine if you:

- are taking other medicines to treat depression (see 'Taking other medicines')
- are taking St. John's Wort, a herbal treatment (Hypericum perforatum).

- have kidney disease
- have had seizures (fits)
- have had mania
- suffer from bipolar disorder
- have eye problems, such as certain kinds of glaucoma (increased pressure in the eye)
- have a history of bleeding disorders (tendency to develop bruises)
- are at risk of low sodium levels
- are currently being treated with another medicine which may cause liver damage
- are taking other medicines containing duloxetine
- have intolerance to some sugars (see end of Section 2)
- are considering stopping ARICLAIM (see section 3)

ARICLAIM may cause a sensation of restlessness or an inability to sit or stand still. You should tell your doctor if this happens to you.

Thoughts of suicide and worsening of your depression or anxiety disorder

If you are depressed and/or have anxiety disorders you can sometimes have thoughts of harming or killing yourself. These may be increased when first starting antidepressants, since these medicines all take time to work, usually about two weeks but sometimes longer.

You may be more likely to think like this if you:

- have previously had thoughts about killing or harming yourself.
- are a young adult. Information from clinical trials has shown an increased risk of suicidal behaviour in adults aged less than 25 years with psychiatric conditions who were treated with an antidepressant.

If you_have thoughts of harming or killing yourself at any time, contact your doctor or go to a hospital straight away.

You may find it helpful to tell a relative or close friend that you are depressed or have an anxiety disorder, and ask them to read this leaflet. You might ask them to tell you if they think your depression or anxiety is getting worse, or if they are worried about changes in your behaviour.

Use in children and adolescents under 18 years of age

ARICLAIM should normally not be used for children and adolescents under 18 years. Also, you should know that patients under 18 have an increased risk of side-effects such as suicide attempt, suicidal thoughts and hostility (predominantly aggression, oppositional behaviour and anger) when they take this class of medicines. Despite this, your doctor may prescribe ARICLAIM for patients under 18 because he/she decides that this is in their best interests. If your doctor has prescribed ARICLAIM for a patient under 18 and you want to discuss this, please go back to your doctor. You should inform your doctor if any of the symptoms listed above develop or worsen when patients under 18 are taking ARICLAIM. Also, the long-term safety effects concerning growth, maturation, and cognitive and behavioural development of ARICLAIM in this age group have not yet been demonstrated.

Taking other medicines

Please tell your doctor or pharmacist if you are taking or have recently taken any other medicines, including medicines obtained without a prescription. The main ingredient of ARICLAIM, duloxetine, is used in other medicines for other conditions:

• depression, anxiety and urinary incontinence

Using more than one of these medicines at the same time should be avoided. Check with your doctor if you are already taking other medicines containing duloxetine.

Your doctor should decide whether you can take ARICLAIM with other medicines. Do not start or stop taking any medicines, including those bought without a prescription and herbal remedies, before checking with your doctor.

You should also tell your doctor if you are taking any of the following:

Monoamine Oxidase Inhibitors (MAOIs): You should not take ARICLAIM if you are taking, or have recently taken within the last 14 days, another antidepressant medicine called a monoamine oxidase

inhibitor (MAOI). Taking a MAOI together with many prescription medicines, including ARICLAIM, can cause serious or even life-threatening side effects. You must wait at least 14 days after you have stopped taking an MAOI before you can take ARICLAIM. Also, you need to wait at least 5 days after you stop taking ARICLAIM before you take a MAOI.

Medicines that cause sleepiness: These include medicines prescribed by your doctor including benzodiazepines, strong painkillers, antipsychotics, phenobarbital, antihistamines.

Medicines that increase the level of serotonin: triptans, tramadol, tryptophan, SSRIs (such as paroxetine and fluoxetine), tricyclics (such as clomipramine, amitriptyline), pethidine, St John's Wort and venlafaxine. These medicines increase the risk of side effects; if you get any unusual symptom taking any of these medicines together with ARICLAIM, you should see your doctor.

Oral anticoagulants: medicines which thin the blood. These medicines might increase the risk of bleeding.

Taking ARICLAIM with food and drink

ARICLAIM may be taken with or without food. Care should be taken if you drink alcohol while you are being treated with ARICLAIM.

Pregnancy and breast-feeding

Ask your doctor or pharmacist for advice before taking any medicine. Tell your doctor if you:

- become pregnant, or you are trying to become pregnant, while you are taking ARICLAIM. You should use ARICLAIM only after discussing the potential benefits and any potential risks to your unborn child with your doctor.
- are breast-feeding. The use of ARICLAIM while breastfeeding is not recommended. You should ask your doctor or pharmacist for advice.

Driving and using machines

ARICLAIM may make you feel sleepy of dizzy. Do not drive or use any tools or machines until you know how ARICLAIM affects you.

Important information about some of the ingredients of ARICLAIM

ARICLAIM contains **sucrose**. If you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this medicinal product.

3. HOW TO TAKE ARICLAIM

Always take ARICLAIM exactly as your doctor has told you. You should check with your doctor or pharmacist if you are not sure.

The usual dose of ARICLAIM is 1 capsule (60 mg) once a day, but your doctor will prescribe the dose that is right for you.

ARICLAIM is for oral use. You should swallow your capsule whole with a drink of water.

To help you remember to take ARICLAIM, you may find it easier to take it at the same times every day.

Talk with your doctor about how long you should keep taking ARICLAIM. Do not stop taking ARICLAIM without talking to your doctor.

If you take more ARICLAIM than you should

Call your doctor or pharmacist immediately if you take more than the amount of ARICLAIM prescribed by your doctor.

If you forget to take ARICLAIM

If you miss a dose, take it as soon as you remember. However, if it is time for your next dose, skip the missed dose and take only a single dose as usual. Do not take a double dose to make up for a forgotten dose. Do not take more than the daily amount of ARICLAIM that has been prescribed for you in one day.

If you stop taking ARICLAIM

DO NOT stop taking your capsules without the advice of your doctor even if you feel better. If your doctor thinks that you no longer need ARICLAIM he or she will ask you to reduce your dose over at least 2 weeks before stopping treatment altogether.

Some patients who stop taking ARICLAIM suddenly have had symptoms such as:

• dizziness, tingling feelings like pins and needles, sleep disturbances (vivid dreams, nightmares, inability to sleep), feeling restless or agitated, feeling anxious, feeling sick (nausea) or being sick (vomiting), tremor (shakiness), headaches, feeling irritable, diarrhoea, excessive sweating or vertigo. These symptoms are usually not serious and disappear within a few days, but if you have symptoms that are troublesome you should ask your doctor for advice.

If you have any further questions on the use of this product, ask your doctor or pharmacist.

4. POSSIBLE SIDE EFFECTS

Like all medicines, ARICLAIM can cause side effects, although not everybody gets them. These effects are normally mild to moderate and often disappear after a few weeks.

Very common side effects (these can affect more than 1 in 10 patients treated)

• feeling sick (nausea), headache, dry mouth, feeling sleepy and dizziness

Common side effects (these can affect from 1 to 10 users in 100 patients treated)

- tiredness, trouble sleeping, anxiety, feeling agitated or having abnormal dreams
- tremor or numbness, including numbness or tingling of the skin
- diarrhoea, constipation, being sick (vomiting), heartburn, breaking wind, stomach pain
- tinnitus (perception of sound in the ear when there is no external sound)
- blurred evesight
- feeling the heart pumping in the chest, flushing, increased sweating, night sweats
- problems getting an erection, less sex drive
- (itchy) rash
- muscle pain, muscle tightness, muscle spasm
- increased yawning
- lack of appetite, weight loss

Uncommon side effects (these can affect from 1 to 10 users in 1,000 patients treated)

- throat inflammation
- feeling disorientated, feeling sleepy, lack of motivation
- tasting things differently than usual, disturbance in attention, stiffness, spasms and involuntary movements of the muscles, muscle twitching, abnormal manner of walking
- poor sleep quality
- restless legs syndrome
- burping, indigestion, gastroenteritis
- vertigo, ear pain

- inflammation of the liver that may cause abdominal pain
- large pupils (the dark centre of the eye), visual disturbance
- fast or irregular heart beat
- sexual problems, including changes in ejaculation and abnormal orgasm
- abnormal periods, including heavy or prolonged periods
- allergic reactions, increased tendency to bruise, blisters or sensitivity to sunlight
- increase in blood pressure, feeling cold in your fingers and/or toes, feeling dizzy (particularly when standing up too quickly), cold sweats, shivering or fainting
- an increased level of sugar in the blood
- need to pass more urine than normal, need to pass urine during the night, difficulty or inability to pass urine or having an urine flow decreased
- grinding of teeth, feeling hot/cold, thirst, throat tightness, nose bleeds
- weight gain

Rare side effects (these can affect from 1 to 10 users in 10,000 patients treated)

- decreased thyroid gland activity
- dehydration
- mania (over activity, racing thoughts and decrease need for sleep)
- bad breath
- increased pressure in the eye (glaucoma)
- menopausal symptoms
- contraction of the jaw muscle
- increased level of cholesterol in the blood, low levels of sodium in the blood (the symptoms are feeling sick and unwell with weak muscles or confused),
- serious allergic reaction which causes difficulty in breathing or dizziness or hives
- fits

Other possible side effects

- hallucinations, suicidal thoughts, behaviour aggression and anger
- a sensation of restlessness or an inability to sit or stand still or "Serotonin syndrome" (a rare reaction which may cause feelings of great happiness, drowsiness, clumsiness, restlessness, feeling of being drunk, fever, sweating or rigid muscles)
- passing bright red blood in your stools, vomiting blood, or black tarry stools (faeces)
- having abnormal urine odour
- syndrome of inadequate secretion of anti-diuretic hormone (SIADH)
- chest pain
- yellow colouration of the skin (jaundice), hepatic failure, Stevens-Johnson syndrome, sudden swelling of skin or mucosa (angioedema)

If any of the side effects gets serious, or if you notice any side effects not listed in this leaflet, please tell your doctor.

5. HOW TO STORE ARICLAIM

Keep out of the reach and sight of children

Do not use ARICLAIM after the expiry date which is stated on the carton.

Store in the original package. Do not store above 30°C.

6. FURTHER INFORMATION

What ARICLAIM contains

The active substance is duloxetine.

Each capsule contains 30 or 60 mg of duloxetine (as hydrochloride).

The **other** ingredients are:

Capsule content: hypromellose, hypromellose acetate succinate, sucrose, sugar spheres, talc, titanium dioxide (E171), triethyl citrate.

(See end of Section 2 for further information on sucrose)

Capsule shell: gelatin, sodium lauryl sulphate, titanium dioxide (E171), indigo carmine (E132), iron oxide yellow (E172) (60 mg only) and edible green ink (30 mg) or edible white ink (60 mg).

Edible green ink: black iron oxide-synthetic (E172), yellow iron oxide-synthetic (E172), propylene glycol, shellac.

Edible White Ink: titanium dioxide (E171), propylene glycol, shellac, povidone.

What ARICLAIM looks like and contents of the pack

ARICLAIM is a hard gastro-resistant capsule.

Each capsule of ARICLAIM contains pellets of duloxetine hydrochloride with a covering to protect them from stomach acid.

ARICLAIM is available in two strengths: 30 and 60 mg.

The 30 mg capsules has an opaque white body, imprinted with '30 mg' and an opaque blue cap, imprinted with '9543'.

The 60 mg capsules has an opaque green body, imprinted with '60 mg' and an opaque blue cap, imprinted with '9542'.

ARICLAIM 30 mg is available in packs of 7, 28 and 98 capsules.

ARICLAIM 60 mg is available in packs of 28 and 98 capsules.

Not all pack sizes may be marketed.

Marketing Authorisation Holder and Manufacturer

Marketing Authorisation Holder: Eli Lilly Nederland BV, Grootslag 1-5,NL-3991 RA Houten, The Netherlands.

Manufacturer: Lilly S.A., Avda. De la Industria, 30,28108 Alcobendas, Madrid, Spain.

For any information about this medicine, please contact the local representative of the Marketing Authorisation Holder:

België/Belgique/Belgien

S.C.S. Boehringer Ingelheim Comm.V.

Tél/Tel: +32 27 73 33 11

България

ТП "Ели Лили Недерланд" Б.В. - България

тел. + 359 2 491 41 40

Česká republika

Boehringer Ingelheim spol. s r.o.

Tel.: + 42 02 34 65 51 11

Danmark

Boehringer Ingelheim Danmark A/S

Tlf: +45 39 15 88 88

Deutschland

Boehringer Ingelheim Pharma GmbH & Co. KG

Tel: +49 (0) 69 50 50 83 09

Eesti

Boehringer Ingelheim Pharma GmbH

Tel: + 37 2 60 80 940

Luxembourg/Luxemburg

S.C.S. Boehringer Ingelheim Comm.V.

Tél/Tel: +32 2 773 33 11

Magyarország

Boehringer Ingelheim Pharma

Tel.: +36 1 224 7120

Malta

Boehringer Ingelheim Ltd.

Tel: +356 25600 500

Nederland

Boehringer Ingelheim b.v.

Tel: +31 30 6 02 59 14

Norge

Boehringer Ingelheim Norway KS

Tlf: +47 66 76 13 00

Österreich

Boehringer Ingelheim Austria GmbH

Tel: +43 1 710 3739

Ελλάδα

ΦΑΡΜΑΣΕΡΒ-ΛΙΛΛΥ Α.Ε.Β.Ε.

Τηλ: +30 210 629 4600

España

Dista S.A..

Tel: + 34 91 623 17 32

France

Boehringer Ingelheim France S.A.S.

Tél: +33 3 26 50 45 33

Ireland

Boehringer Ingelheim Ireland Ltd.

Tel: +353-(0) 1 661 4377

Ísland

Eli Lilly Danmark A/S, Útibú á Íslandi

Tel: + 354 520 34 00

Italia

Eli Lilly Italia S.p.A. Tel: + 39- 055 42571

161: + 39- 055 425

Κύπρος

Boehringer Ingelheim Ellas A.E.

Τηλ: +30 2 10 89 06 300

Latvija

Boehringer Ingelheim Pharma GmbH

Tel: +37 167 24 00 68

Lietuva

Boehringer Ingelheim Pharma Ges mbH

Tel.: +370 37 47 39 22

Polska

Boehringer Ingelheim Sp.z o.o.

Tel.: +48 22 699 0 699

Portugal

Boehringer Ingelheim, Lda

Tel: +351 21 313 53 00

România

Eli Lilly România S.R.L.

Tel: + 40 21 4023000

Slovenija

Boehringer Ingelheim Pharma

Tel.: +386 1 586 40 00

Slovenská republika

Boehringer Ingelheim Pharma

Tel.: +421 2 5341 8445

Suomi/Finland

Oy Eli Lilly Finland Ab

Puh/Tel: +358 9 8545 250

Sverige

Boehringer Ingelheim AB

Tel: +46 8 721 21 00

United Kingdom

Boehringer Ingelheim Ltd.

Tel: +44 (0) 1256 315999

This leaflet was last approved in

Detailed information on this medicine is available on the European Medicines Agency (EMEA) web site: http://www.emea.europa.eu.